FILE 'HOME' ENTERED AT 07:02:33 ON 25 SEP 2002

=> file reg COST IN U.S. DOLLARS SINCE FILE TOTAL SESSION 0.21 0.21

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 23 SEP 2002 HIGHEST RN 454421-17-1 DICTIONARY FILE UPDATES: 23 SEP 2002 HIGHEST RN 454421-17-1

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

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Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

*** YOU HAVE NEW MAIL ***

=> Uploading 09880727.str

L1 STRUCTURE UPLOADED

=> s 1 full <----->user Break-----> u SEARCH ENDED BY USER

=> d l1 L1 HAS NO ANSWERS

Structure attributes must be viewed using STN Express query preparation.

=> s l1 full FULL SEARCH INITIATED 07:03:30 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 2487 TO ITERATE

100.0% PROCESSED 2487 ITERATIONS

535 ANSWERS

SEARCH TIME: 00.00.01

L2 535 SEA SSS FUL L1

=> file caplus COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 140.28 140.49

FULL ESTIMATED COST

FILE 'CAPLUS' ENTERED AT 07:03:37 ON 25 SEP 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 25 Sep 2002 VOL 137 ISS 13 FILE LAST UPDATED: 23 Sep 2002 (20020923/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For

information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

=> s 12 703 L2 L3

=> s 13 and label 48825 LABEL

3 L3 AND LABEL T.4

=> s 13 and label? 374492 LABEL? 36 L3 AND LABEL? 1.5

=> d 14 bib abs hitstr 1-3

ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS T.4

1981:186601 CAPLUS AN

DN 94:186601

DNA repair in pollen: range of mutagens inducing repair, effect of TТ replication inhibitors and changes in thymidine nucleotide metabolism during repair

Jackson, J. F.; Linskens, H. F. ΑU

Waite Agric. Res. Inst., Univ. Adelaide, Glen Osmond, 5064, Australia CS

MGG, Mol. Gen. Genet. (1980), 180(3), 517-22 SO CODEN: MGGEAE; ISSN: 0026-8925

DTJournal

English LΑ

AB

Pollen of Petunia hybrida carry out DNA repair during the 1st 2 h of germination when certain mutagens are included in the germination medium. This repair, detected as unscheduled DNA synthesis, since there is no replicative DNA synthesis in Petunia pollen, can be induced by the chem. mutagens, N-methyl-N'-nitro-N-nitrosoguanidine [70-25-7], 4-nitroquinoline 1-oxide [56-57-5], azaserine [115-02-6], and methyl methanesulfonate [66-27-3]. These compds. are all capable of direct covalent interaction with DNA. Mutagens requiring metabolic activation before interaction with DNA did not induce DNA repair synthesis in pollen. The practice of solubilizing water-insol. chem. mutagens with DMSO [67-68-5] did not prove practical, due to the extremely harmful effects of DMSO on pollen. Pretreatment of pollen before germination with pure Et20 [60-29-7], however, had no harmful effect on either repair or pollen germination. Therefore, water-insol., Et20-sol. mutagens were tested by pretreatment of the pollen with mutagens in Et20 soln. By this means, the direct-acting mutagen, Et2SO4 [64-67-5], also brings about unscheduled DNA synthesis in pollen, while 2-acetylaminofluorene [53-96-3] and dimethyl-p-aminobenzene [60-11-7], both requiring metabolic activation, did not do so. Inhibitors of DNA replicative synthesis, hydroxyurea [127-07-1], azaserine, azauridine [54-25-1], and fluorodeoxyuridine [50-91-9] did not inhibit unscheduled DNA synthesis brought about by N-methyl-N'-nitro-N-nitrosoguanidine. On the contrary, these compds. stimulated repair synthesis to varying degrees, hydroxyurea having the greatest effect. Pollen uptake of 3H-labeled thymidine [50-89-5] and the amt. of radioactive label subsequently appearing in dTMP and dTDP + dTTP was increased by 4-nitroquinoline 1-oxide. Partial inhibition of these increases and of 4-nitroquinoline 1-oxide induced repair synthesis by cAMP [60-92-4] suggested that thymidine: AMP phosphotransferase [60440-28-0] rather than thymidine kinase was responsible for thymidine phosphorylation in pollen. Enzyme assays on pollen exts. confirmed this.

IT 54-25-1

RL: BIOL (Biological study)

(DNA repair by Petunia hybrida pollen response to)

54-25-1 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA

INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS

AN 1980:3705 CAPLUS

DN 92:3705

TI Simultaneous estimation of rates of pyrimidine and purine nucleotide synthesis de novo in cultured human cells

AU Huisman, William H.; Raivio, Kari O.; Becker, Michael A.

CS Rheumatol. Sect., VA Hosp., San Diego, CA, 92161, USA

SO J. Biol. Chem. (1979), 254(24), 12595-602 CODEN: JBCHA3; ISSN: 0021-9258

DT Journal

LA English

AΒ

The requirement of the pathways of pyrimidine and purine nucleotide synthesis de novo for CO2 was exploited in a method for simultaneous estn. of the rates of operation of these pathways in cultured human lymphoblasts and fibroblasts. Rates of incorporation of H14CO3- into pyrimidine and purine compds. were const. for .ltoreq.2 h and were proportional to the cell no. in the assay. Incorporation rates appeared to reflect the rates of synthesis of pyrimidine and purine compds. in individual cell strains under conditions in which: (1) carbamyl phosphate concns. were <0.5% of the hourly flux of H14CO3- into the pyrimidine pathway; and (2) the sp. activities of HCO3- pools were apparently unchanged. Alterations in the rates of H14CO3- incorporation during incubation of normal and hypoxanthine-guanine phosphoribosyltransferase-deficient lymphoblasts with purine bases and purine and pyrimidine nucleosides were in agreement with previous observations using alternative methods for the individual estn. of rates of purine or pyrimidine synthesis. In addn., comparable increases in H14CO3- incorporation into purines and pyrimidines were demonstrated in human lymphocytes during exposure to phytohemagglutinin, a stimulus previously shown to accelerate rates of pyrimidine and purine synthesis. These findings provided evidence for the validity of the present method in assessing the rates of pyrimidine and purine synthesis. High correlations between log phase growth rates of individual lymphoblast lines and their rates of incorporation of label were obsd. Although specific and consistent differences were obsd. in the rates of H14CO3- incorporation into pyrimidine and purine compds. in normal, hypoxanthine-guanine phosphoribosyltransferase-deficient, and 5-phosphoribosyl 1-pyrophosphate synthetase superactive strains, lack of information concerning the sp. activities of intracellular HCO3- pools in different cell strains restricted abs. comparisons of the rates of nucleotide synthesis between cell strains.

IT 54-25-1

RL: BIOL (Biological study)

(purine and pyrimidine nucleotide formation from bicarbonate by fibroblast and lymphoblast in response to)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS L4

1974:491480 CAPLUS AN

81:91480 DN

Preparation of 6-azauracil-5-t and 6-azauridine-5-t of high molar activity TI

Filip, Jiri; Skoda, Jan; Hradec, Hynek ΑU

Inst. Res. Prod. Uses Radioisot., Czech. Acad. Sci., Prague, Czech. CS

J. Label. Compounds (1974), 10(1), 59-71 SO CODEN: JLCAAI

Journal \mathbf{DT}

English LA

Catalytic reductive dehalogenation of 5-bromo-6-azauracil with carrier-free T gave 6-azauracil-5-3H of a molar activity of 19.0Ci/mmole. AΒ The conditions for the catalytic reductive dehalo-genation were examd. in tracer expts. Microbial transformation gave 6-azauridine-5-3H from 6-azauracil-5-3H having molar activity of 18.8 Ci/mmole. The stability of T was investigated in aq. medium at 100.degree.C.

53615-16-0P ITRL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and label stability of)

53615-16-0 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione-6-t, 2-.beta.-D-ribofuranosyl- (9CI) (CA CNINDEX NAME)

Absolute stereochemistry.

=> s 15 not 14

33 L5 NOT L4

=> d 16 bib abs hitstr 1-33

ANSWER 1 OF 33 CAPLUS COPYRIGHT 2002 ACS

2001:851808 CAPLUS ΑN

135:367666 DN Nucleotide analogs and their use in labeling nucleic acids for TIhybridization assays

McGall, Glenn; Barone, Anthony D. IN

PΑ USA SO U.S. Pat. Appl. Publ., 47 pp., Cont.-in-part of U.S. Appl. 2001 18,514.

CODEN: USXXCO

DT Patent LA English

FAN.CNT 2

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | | |
|---------------------|------|----------|-----------------|----------|--|--|
| | | | | | | |
| PI US 2001044531 | A1 | 20011122 | US 2001-780574 | 20010209 | | |
| US 2001018514 | A1 | 20010830 | US 1998-126645 | 19980731 | | |
| PRAI US 1998-126645 | A2 | 19980731 | | | | |

OS MARPAT 135:367666

AB Nucleic acid labeling compds. contg. heterocyclic derivs. are disclosed. The heterocyclic deriv. contg. compds. are synthesized by condensing a heterocyclic deriv. with a cyclic group (e.g. a ribofuranose deriv.). The labeling compds. are suitable for enzymic attachment to a nucleic acid, either terminally or internally, to provide a mechanism of nucleic acid detection. Thus, a no. of biotin- or fluorescein purine- and pyrimidine-.beta.-D-ribofuranoside analogs were prepd. These analogs were successfully incorporated into hybridization probes (using terminal deoxynucleotidyltransferase) and utilized in single nucleotide polymorphism geno-typing using micro-chip arrays.

IT 257297-94-2P 257298-04-7P
 RL: BSU (Biological study, unclassified); IMF (Industrial manufacture);
 SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
 (nucleotide analogs and their use in labeling nucleic acids
 for hybridization assays)

RN 257297-94-2 CAPLUS

CN 1H-Thieno[3,4-d]imidazole-4-pentanamide, hexahydro-2-oxo-N-[6-oxo-6-[[3-[2,3,4,5-tetrahydro-2-[5-0-[hydroxy[[hydroxy(phosphonooxy)phosphinyl]oxy]phosphinyl]-.beta.-D-ribofuranosyl]-3,5-dioxo-1,2,4-triazin-6-yl]-2-propenyl]amino]hexyl]-, (3aS,4S,6aR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

PAGE 1-B

RN 257298-04-7 CAPLUS

CN Spiro[isobenzofuran-1(3H),9'-[9H]xanthene]-5-carboxamide,

3',6'-dihydroxy-3-oxo-N-[6-oxo-6-[[3-[2,3,4,5-tetrahydro-2-[5-0-[hydroxy[[hydroxy(phosphonooxy)phosphinyl]oxy]phosphinyl]-.beta.-Dribofuranosyl]-3,5-dioxo-1,2,4-triazin-6-yl]-2-propenyl]amino]hexyl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

> PAGE 1-A HO_

HO OPO3H2 HO OH S R R R N N N (CH2) 5
$$\frac{H}{H}$$

PAGE 1-B

257297-91-9P 257297-92-0P 257297-93-1P ITRL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (nucleotide analogs and their use in labeling nucleic acids for hybridization assays)

257297-91-9 CAPLUS RN

1H-Isoindole-1,3(2H)-dione, 2-[3-[2,3,4,5-tetrahydro-3,5-dioxo-2-(2,3,5-CN tri-O-benzoyl-.beta.-D-ribofuranosyl)-1,2,4-triazin-6-yl]-2-propenyl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

RN 257297-92-0 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 6-(3-amino-1-propenyl)-2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

RN 257297-93-1 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N-[3-(2,3,4,5-tetrahydro-3,5-dioxo-2-.beta.-D-ribofuranosyl-1,2,4-triazin-6-yl)-2-propenyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

IT 2169-64-4

RL: RCT (Reactant); RACT (Reactant or reagent) (nucleotide analogs and their use in labeling nucleic acids for hybridization assays)

RN 2169-64-4 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-(2,3,5-tri-O-acetyl-.beta.-D-ribofuranosyl)- (9CI) (CA INDEX NAME)

ANSWER 2 OF 33 CAPLUS COPYRIGHT 2002 ACS 1.6

2000:758718 CAPLUS AN

DN 135:40384

Structure-inhibitory profiles of nucleosides for the human intestinal N1 ΤI and N2 Na+-nucleoside transporters

Patil, Shivakumar D.; Ngo, Leock Y.; Unadkat, Jashvant D. ΑU

H272 Health Sciences, Department of Pharmaceutics, University of CS Washington, Seattle, WA, 98195, USA

Cancer Chemotherapy and Pharmacology (2000), 46(5), 394-402 SO CODEN: CCPHDZ; ISSN: 0344-5704

PΒ Springer-Verlag

Journal DT

LΑ English

The structure-inhibitory profiles of nucleosides for the N1 and N2 AB Na+-nucleoside transporters of the human intestine were detd. The uptake of 3H-labeled prototypic substrates of the N1 (inosine) and N2 (thymidine) transporters into human intestinal brush border membrane vesicles was measured by a rapid filtration technique in the presence and absence of various uridine and adenosine analogs and antiviral and anticancer nucleoside drugs (100 and 1000 .mu.M). In the ribose ring, the 3'-oxygen is required for inhibition of uptake of nucleosides by both the N1 and N2 transporters. The structural requirements for such inhibition differ with respect to modifications on the 5' position of the sugar ring or on the base. The N2 transporter is more tolerant to these substitutions than is the N1 transporter. The 6 position on uracil and the 8 position on adenine are crit. for inhibition of uptake of nucleosides by both the N1 and N2 nucleoside transporters. These data are the 1st evidence that the binding site(s) of the human N1 and N2 transporters differ in their interaction with analogs of their common substrates, uridine and adenosine. Such studies can provide insight into the crit. structural determinants of the substrate necessary for recognition by the Na+-nucleoside transporters of the human intestine.

TΤ **54-25-1**, 6-Azauridine RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(structure-inhibitory profiles of nucleosides for human intestinal N1 and N2 Na+-nucleoside transporters)

RN54-25-1 CAPLUS

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA CN INDEX NAME)

RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L6
     ANSWER 3 OF 33 CAPLUS COPYRIGHT 2002 ACS
AN
     2000:98825 CAPLUS
DN
     132:133201
ΤI
     Nucleotide analogs and their use in labeling nucleic acids for
     hybridization assays
IN
     McGall, Glenn H.; Barone, Anthony D.
PA
     Affymetrix, Inc., USA
SO
     PCT Int. Appl., 69 pp.
     CODEN: PIXXD2
DТ
     Patent
LA
     English
FAN.CNT 2
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
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PΙ
     WO 2000006771
                     A2
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                                          WO 1999-US12390 19990720
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    EP 1124838
                      A2 . 20010822
                                          EP 1999-937150
                                                            19990720
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             IE, SI, LT, LV, FI, RO
     JP 2002521495
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                           20020716
                                          JP 2000-562553
                                                            19990720
PRAI US 1998-126645
                      Α
                           19980731
    WO 1999-US12390
                      W
                            19990720
OS
    MARPAT 132:133201
    Nucleic acid labeling compds. contg. heterocyclic derivs. are
    disclosed. The heterocyclic deriv. contg. compds. are synthesized by
    condensing a heterocyclic deriv. with a cyclic group (e.g. a ribofuranose
    deriv.). The labeling compds. are suitable for enzymic
    attachment to a nucleic acid, either terminally or internally, to provide
    a mechanism of nucleic acid detection. Thus, a no. of biotin- or
    fluorescein purine- and pyrimidine-.beta.-D-ribofuranoside analogs were
            These analogs were successfully incorporated into hybridization
    probes (using terminal deoxynucleotidyltransferase) and utilized in single
    nucleotide polymorphism genotyping using microchip arrays.
IT
    257297-94-2P 257298-04-7P
    RL: BPR (Biological process); BSU (Biological study, unclassified); SPN
```

(nucleotide analogs and their use in **labeling** nucleic acids for hybridization assays)

(Synthetic preparation); BIOL (Biological study); PREP (Preparation); PROC

RN 257297-94-2 CAPLUS

CN 1H-Thieno[3,4-d]imidazole-4-pentanamide, hexahydro-2-oxo-N-[6-oxo-6-[[3-[2,3,4,5-tetrahydro-2-[5-O-[hydroxy[[hydroxy(phosphonooxy)phosphinyl]oxy]phosphinyl]-.beta.-D-ribofuranosyl]-3,5-dioxo-1,2,4-triazin-6-yl]-2-propenyl]amino]hexyl]-, (3aS,4S,6aR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

PAGE 1-B

RN 257298-04-7 CAPLUS

CN Spiro[isobenzofuran-1(3H),9'-[9H]xanthene]-5-carboxamide,
3',6'-dihydroxy-3-oxo-N-[6-oxo-6-[[3-[2,3,4,5-tetrahydro-2-[5-0-[hydroxy[[hydroxy(phosphonooxy)phosphinyl]oxy]phosphinyl]-.beta.-D-ribofuranosyl]-3,5-dioxo-1,2,4-triazin-6-yl]-2-propenyl]amino]hexyl](9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

PAGE 1-A

HO OPO3H2 HO OH N O (CH2)
$$\frac{H}{H}$$

IT 2169-64-4

RL: RCT (Reactant); RACT (Reactant or reagent) (nucleotide analogs and their use in labeling nucleic acids for hybridization assays)

RN 2169-64-4 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-(2,3,5-tri-O-acetyl-.beta.-D-ribofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 257297-91-9P 257297-92-0P 257297-93-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(nucleotide analogs and their use in **labeling** nucleic acids for hybridization assays)

RN 257297-91-9 CAPLUS

CN 1H-Isoindole-1,3(2H)-dione, 2-[3-[2,3,4,5-tetrahydro-3,5-dioxo-2-(2,3,5-tri-O-benzoyl-.beta.-D-ribofuranosyl)-1,2,4-triazin-6-yl]-2-propenyl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

RN 257297-92-0 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 6-(3-amino-1-propenyl)-2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

RN 257297-93-1 CAPLUS

CN Acetamide, 2,2,2-trifluoro-N-[3-(2,3,4,5-tetrahydro-3,5-dioxo-2-.beta.-D-ribofuranosyl-1,2,4-triazin-6-yl)-2-propenyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry unknown.

L6 ANSWER 4 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1998:759665 CAPLUS

DN 130:95775

TI Synthesis of tritium-labeled diazines and their analogs

AU Myasoedov, Nykolai F.; Sidorov, Georgy V.

CS Institute of Molecular Genetics, RAS, Moscow, 123182, Russia

SO Journal of Labelled Compounds & Radiopharmaceuticals (1998), 41(11), 993-1003

CODEN: JLCRD4; ISSN: 0362-4803

PB John Wiley & Sons Ltd.

DT Journal

LA English

AB Some 40 diazines have been tritiated to high specific activities using a variety of labeling procedures such as catalytic hydrogen isotope exchange both in soln. and the solid state, redn. and hydration. For purine derivs. it is shown that the solid state catalytic isotope exchange reaction is the most effective method. With pyrimidines this reaction is accompanied by a parallel hydration reaction of the 5,6-double bond to form a complex mixt. of products. Identification and quant. estn. of these products has been accomplished in terms of the reaction condition (solvent, nature of catalyst). Key Words: tritium, catalytic hydrogenation, purines, pyrimidines, nucleosides, nucleotides, phytohormones, and terminators of DNA synthesis.

IT 219524-90-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of using solid-state isotope exchange reaction)

RN 219524-90-0 CAPLUS

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl-, labeled with CN tritium (9CI) (CA INDEX NAME)

Absolute stereochemistry.

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 9 ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L6
     ANSWER 5 OF 33 CAPLUS COPYRIGHT 2002 ACS
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1994:331137 CAPLUS AN

DN 120:331137

Pharmaceutical liposomes comprising lipids derivatized with PEG for ΤI treatment of inflamed tissues

IN Woodle, Martin C.; Martin, Francis J.; Huang, Shi Kun

Liposome Technology, Inc., USA PA

PCT Int. Appl., 97 pp. SO

CODEN: PIXXD2

DT Patent

LAEnglish

| FAN. | CNT | 9 | | | | | | | | | | | | |
|------|-----|-------------|--------|-----------|-----|---------|-------|-------|-----|------|------|-----|-----|----|
| | PAT | TENT NO. | KIND | DATE | | APPLI | CATI | ON NO | ο. | DATE | | | | |
| | | | | | | | | | | | | | | |
| PI | WO | 9407466 | A1 | 19940414 | | WO 19 | 93-U | S9572 | 2 | 1993 | 1007 | | | |
| | | W: AU, CA, | JP | | | | | | | | | | | |
| | | RW: AT, BE, | CH, DE | , DK, ES, | FR, | GB, GR, | ΙE, | ΙΤ, | LU, | MC, | NL, | PT, | SE | |
| | US | 5356633 | A | 19941018 | | US 19 | 92-9! | 58100 | 0 | 1992 | 1007 | | | |
| | EΡ | 662820 | A1 | 19950719 | | EP 19 | 93-92 | 2329 | 5 | 1993 | 1007 | | | |
| | ΕP | 662820 | B1 | 19970507 | | | | | | | | | | |
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| PRAI | US | 1992-958100 | A | 19921007 | | | | | | | | | | |
| | US | 1989-425224 | A2 | 19891020 | | | | | | | | | | |
| | US | 1991-642321 | A2 | 19910115 | | | | | | • | | | | |
| | WO | 1993-US9572 | W | 19931007 | | | | | | | | | | |

AΒ Pharmaceutical liposomes comprising vesicle-forming lipids derivatized with PEG for delivery to an inflamed region are disclosed. After i.v. administration, the liposomes are taken up by the inflamed region within 24-48 h, for site-specific release of the therapeutic compd. into the inflamed region. PEG conjugates with distearylphosphatidylethanolamine (prepn. given) was combined with partially hydrogenated egg phosphatidylcholine in a ratio of 0.1:2 and the lipid mixt. was hydrated and extruded through a 0.1 .mu.m polycarboante membrane to produce multilamellar vesicle with av. size .apprx.0.1.mu.m. Above liposomes were labeled and injected to mice and the concn. of liposomes in the blood was detd. 24 after injection. The amt. of dose remaining in the blood 24 h after injection was 5-40% as compared to <1% for liposomes lacking PEG-derivatized lipids. Extravasation of liposomes contg. PEG-derivatized lipids into sites of bradykinin-induced inflammation in rats was studied.

IT**2169-64-4**, Azaribine

RL: BIOL (Biological study)

(pharmaceutical liposomes comprising lipids derivatized with PEG and, for treatment of inflamed tissues)

2169-64-4 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-(2,3,5-tri-O-acetyl-.beta.-D-CNribofuranosyl) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

1.6 ANSWER 6 OF 33 CAPLUS COPYRIGHT 2002 ACS

1994:280292 CAPLUS ΑN

DN 120:280292

ΤI Detection and therapy of lesions with biotin/avidin conjugates

Goldenberg, David M. IN

PA Immunomedics, Inc., USA

SO PCT Int. Appl., 62 pp.

CODEN: PIXXD2

DTPatent

English

IT

| FAN. | CNT | 14 | | | | | | | | |
|------|------|--------------|--------|-----------|-----|--------------------|-----------|-----|-----|----|
| | PAT | TENT NO. | KIND | DATE | | APPLICATION NO. | DATE | | | |
| | | | | | | | | | | |
| ΡI | WO | 9404702 | A2 | 19940303 | | WO 1993-US7754 | 19930820 | | | |
| | WO | 9404702 | A3 | 19961003 | | | | | | |
| | | W: AU, CA, | JP | | | | | | | |
| | | RW: AT, BE, | CH, DE | , DK, ES, | FR, | GB, GR, IE, IT, LU | , MC, NL, | PT, | SE | |
| | US | 5525338 | Α | 19960611 | | US 1992-933982 | 19920821 | | | |
| | ΕP | 656115 | A1 | 19950607 | | EP 1993-920155 | 19930820 | | | |
| | | R: AT, BE, | CH, DE | , DK, ES, | FR, | GB, GR, IE, IT, LI | , LU, MC, | NL, | PT, | SE |
| | ΑU | 671489 | B2 | 19960829 | | AU 1993-50184 | 19930820 | | | |
| | JP | 09505799 | T2 | 19970610 | | JP 1993-506499 | 19930820 | | | |
| PRAI | US | 1992-933982 | Α | 19920821 | | | | | | |
| | 1.10 | 1000 1107764 | 7.7 | 1000000 | | | | | | |

19930820 WO 1993-US7754 W Lesions in a patient are targeted for detection or therapy by parenteral AB injection of (A) biotin or avidin conjugated to a protein which binds to a marker substance produced by or assocd. with the lesion; (B) optionally, a clearing compn. comprising avidin (after a biotin-protein conjugate) or biotin (after an avidin-protein conjugate) to clear compn. (A) from nontargeted sites; (C) a detection or therapeutic compn. comprising a conjugate of avidin or biotin, binding protein as in (A), and a detection or therapeutic agent; (D) optionally, a conjugate of avidin or biotin with a detection or therapeutic agent. The lesion may be cancerous, cardiovascular (e.g. thrombus, embolus, infarct, atherosclerotic plaque), infectious, or inflammatory. The binding protein may be a hormone, lymphokine, growth factor, enzyme, immunomodulator, receptor, (monoclonal) antibody (fragment), etc. The detection agent may be a radionuclide, MRI enhancing agent, photoactivated dye, etc. The therapeutic agent may be an isotope, drug, toxin, hormone, receptor antagonist, etc. Kits contg. sterile injectable compns. for use with embodiments of this method are described. Thus, a patient with a colon neoplasm was injected i.v. with a biotinylated monoclonal antibody to CEA. Two days later, unlabeled avidin was injected i.v., followed the next day by biotinylated antibody to colon-specific antigen p labeled with 111In. Scanning with a .gamma. camera 2 days later revealed radioactivity in a lesion in the sigmoid colon, in agreement with sigmoidoscopy findings.

2169-64-4D, Azaribine, conjugates with avidin or biotin and

binding protein

RL: BIOL (Biological study)

(lesion targeting with, for detection and therapy)

RN 2169-64-4 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-(2,3,5-tri-O-acetyl-.beta.-D-ribofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 7 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1989:72943 CAPLUS

DN 110:72943

TI Uracil ribonucleotide metabolism in rat and human glomerular epithelial and mesangial cells

AU Dumler, Francis; Cortes, Pedro

CS Dep. Med., Henry Ford Hosp., Detroit, MI, 48202, USA

SO Am. J. Physiol. (1988), 255(6, Pt. 1), C712-C718 CODEN: AJPHAP; ISSN: 0002-9513

DT Journal

LA English

Culture of rat mesangial cells in medium contg. dialyzed fetal calf serum resulted in UTP loss (28 nmol/mg DNA/h); the addn. of 2 .mu.M orotate to this medium resulted in net UTP accretion (5.42 nmol/mg DNA/h). Rat mesangial cells demonstrated 16- and 29-46-fold greater UTP and UDP-sugar pools, resp., than whole glomeruli. In human mesangial cells, 6-azauridine (500 .mu.M) decreased UDP-sugar pools by 48%, whereas uridine (50 .mu.M) produced a 2.5-fold increase. Human and rat mesangial cells had greater (1.8-6.1-fold) UDP-sugar pools than epithelial cells and 1.7-3.4-fold greater labeled precursor incorporation into EDP-sugars. Thus, glomerular cells utilize both exogenous orotate and uridine for ribonucleotide synthesis, and the extracellular concn. of these precursors markedly influence the formation and cellular content of DUP-sugars. This may represent diverse activity of glycosylating reactions.

IT **54-25-1**, 6-Azauridine

RL: BIOL (Biological study)

(uracil ribonucleotide metabolic pools response to, in human glomerular mesangial cells)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

L6 ANSWER 8 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1988:33029 CAPLUS

DN 108:33029

TI Cloning and expression of the OMP decarboxylase gene URA4 from Schizosaccharomyces pombe

AU Bach, Marie Louise

CS Lab. Genet. Physiol., IBMC, Strasbourg, F-67084, Fr.

SO Curr. Genet. (1987), 12(7), 527-34

CODEN: CUGED5; ISSN: 0172-8083 DT Journal

LA English

AB URA4, The gene coding for orotidine monophosphate decarboxylaase (OMPdecase), was cloned from the fission yeast by homologous complementation and restricted in an Escherichia coli-S. pombe replicative plasmid to a 1.76-kb HindII fragment. This plasmid is maintained at a high copy no. in S. pombe and allows OMPdecase expression in Saccharomyces cerevisiae as well as in E. coli. After characterization by restriction mapping and Southern hybridization, the cloned gene was used as a probe to measure URA4 transcription and to examine its regulation. MRNA levels were measured by DNA/RNA filter-hybridization with pulse-labeled RNAs during 6-azauridine (6-AUR)-inhibited growth in wild-type and 6-AUR-sensitive strains. In S. pombe, the OMP analog 6-AUR does not regulate the level of OMPdecase formation as it does in S. cerevisiae, but rather modifies the ratio of total polyA+ to polyA- RNAs in the cell. Based on these results and on corresponding enzyme activities, this study demonstrates divergent pyrimidine pathway regulation in the 2 yeasts S. cerevisiae and S. pombe. Finally, the use of the URA4 gene as a convenient selective marker for genetic engineering in S. pombe is proposed.

IT **54-25-1**, 6-Azauridine

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(orotidine monophosphate decarboxylase gene of Schizosaccharomyces pombe regulation by, in Saccharomyces cerevisiae)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

L6 ANSWER 9 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1985:75049 CAPLUS

DN 102:75049

TI Quantitative determination of nucleosides and their phosphate esters. 1. The acidic nucleosides, 3-deazauridine and 6-azauridine

AU Welch, A. D.; Nemec, J.; Panahi, J.

CS Clin. Pharmacol., St. Jude Child. Res. Hosp., Memphis, TN, 38101, USA

Int. J. Biochem. (1984), 16(6), 587-91
CODEN: IJBOBV; ISSN: 0020-711X

DT Journal

SO

LA English

Acidic ribonucleosides, e.g., 3-deazauridine and 6-azauridine, were quant. sepd. from their metab. phoshphorylated esters by chromatog. on minicolumns contg. .apprx.1.8 mL of DEAE-cellulose equilibrated with 10 mM Na phosphate, pH 6.0-6.2. The chem. stable, 3H-labeled nucleosides were eluted from the minicolumns with 10 mM Na phosphate (pH 6.0); subsequently, the nucleotides were eluted completely with 0.5M HCl-0.5M NaCl. Quantitation was based on liq. scintillation counting. For example, this method was used to study phosphorylation of [5-3H]-6-azauridine and recoveries of total radioactivities after incubation with fractions of cultured human RPMI-6410 cells contg. uridine-cytidine kinase activity. [5-3H]-6-azauridine, at levels ranging 6.25 .mu.m-12.8 mM, yielded from 16.4-80.5% phosphorylation; recoveries of radioactivities were .apprx.100%.

IT 53615-16-0

RL: ANT (Analyte); ANST (Analytical study) (detn. of, by DEAE-cellulose chromatog. and liq. scintillation counting)

RN 53615-16-0 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione-6-t, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 54-25-1 2018-19-1

RL: ANT (Analyte); ANST (Analytical study) (detn. of, by chromatog. on DEAE-cellulose minicolumns)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

RN 2018-19-1 CAPLUS CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-(5-O-phosphono-.beta.-D-ribofuranosyl)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 10 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1983:242 CAPLUS

DN 98:242

TI Stimulation by 6-azauridine of carbamoyl phosphate synthesis for pyrimidine biosynthesis in mouse spleen slices

AU Tatibana, Masamiti; Kita, Kazuko; Asai, Takashi

CS Sch. Med., Chiba Univ., Chiba, 280, Japan

SO Eur. J. Biochem. (1982), 128(2-3), 625-9 CODEN: EJBCAI; ISSN: 0014-2956

DT Journal

LA English

GΙ

Slices of spleen from anemic mice were incubated with [14C]bicarbonate in the presence and absence of 6-azauridine (I) [54-25-1] and the amts. of 14C that entered the de novo pyrimidine [289-95-2] biosynthetic pathway were assessed and compared. Compds. analyzed included carbamoylaspartate [13184-27-5], dihydroorotate [155-54-4], orotate [65-86-1] plus its derivs., acid-sol. uracil [66-22-8] and cytosine 5'-nucleotides, nucleic acid pyrimidines, free pyrimidine bases, and nucleosides. As the intracellular levels of carbamoyl phosphate [590-55-6] and acid-sol. deoxyribonucleotides are known to be relatively low, the radioactivities of these compds. were not measured. Degrdn. of labeled uridine was limited in this tissue, therefore the radioactivity of degradative products of pyrimidines was not considered. When the slices were incubated with 0.5 mM 6-azauridine for 10 min and then with [14C]bicarbonate for an addnl. 10 min and 30 min, the sum of

radioactivity found in the above compds., which represents the total amt. of 14C that entered the pyrimidine pathway, was 2.1 and 2.3 times greater than when the tissue slices were incubated in the absence of the analog. When the 14C distribution among the C atoms of the mols. of labeled carbamoylaspartate and uracil was investigated, >90% of the total 14C in these compds. was found to be derived directly from carbamoyl phosphate and the remaining portion was from aspartate, either in the presence or absence of 6-azauridine. There was no indication that 6-azauridine altered [14C]bicarbonate permeation through the cell membrane or its intracellular metab. These results, along with the pattern of early intermediate accumulation seen in the presence of 6-azauridine, indicate that 6-azauridine stimulates the prodn. of carbamoyl phosphate for the pyrimidine biosynthetic pathway in the mouse spleen. Of the radioactive early intermediates which accumulated, only orotate, its derivs. (orotidine [314-50-1] and orotidine 5'-monophosphate [2149-82-8]) or both appeared in the medium, presumably the result of leakage through the cell membranes. Stimulation of the pyrimidine pathway was not obsd. in the case of Ehrlich ascites tumor cells incubated under similar conditions with 6-azauridine.

IT 54-25-1

RL: BIOL (Biological study) (carbamoyl phosphate and pyrimidine biosynthesis response to, in spleen)

54-25-1 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) CNINDEX NAME)

Absolute stereochemistry.

ANSWER 11 OF 33 CAPLUS COPYRIGHT 2002 ACS L6

1981:250 CAPLUS AN

94:250 DN

Selective enhancement of 5-fluorouridine uptake and action in rat ΤI hepatomas in vivo following pretreatment with D-galactosamine and 6-azauridine or N-(phosphonacetyl)-L-aspartate

Anukarahanonta, T.; Holstege, A.; Keppler, D. O. R. ΑU

Biochem. Inst., Univ. Freiburg, Freiburg, D-7800, Fed. Rep. Ger. Eur. J. Cancer (1980), 16(9), 1171-80 CODEN: EJCAAH; ISSN: 0014-2964 CS

SO

DTJournal

English LA

AB The sequential combination of three antipyrimidines was studied in rats carrying Morris hepatoma 7777, the AS-30D ascites hepatoma, or the solid intrahepatic tumor. The uptake of [14C]5-fluorouridine (I) [316-46-1] and its incorporation into RNA was selectively enhanced in hepatomas in vivo by pretreatment of the animals with D-galactosamine [7535-00-4] and an inhibitor of de novo pyrimidine synthesis, such as 6-azauridine **54-25-1**] or N-(phosphonacetyl)-L-aspartate [51321-79-0]. This pretreatment resulted in a transient depletion of uridine phosphate pools which was the prerequisite for the subsequent increase in 5-fluorouridine phosphorylation in the hepatoma. It was demonstrated by radio-paper chromatog. that the formation of 5-fluorouridine diphosphate N-acetylhexosamines was markedly enhanced when the pretreatment included D-galactosamine. The incorporation of labeled precursors into nucleic acids of AS-30D cells treated with 5-fluorouridine indicated that a severe inhibition of thymidylate synthase was assocd. with only a moderate depression of DNA synthesis, as measured by incorporation into DNA of [3H]deoxyuridine and [14C] deoxyadenosine, resp. Survival of rats bearing the intrahepatic or the ascites form of the AS-30D hepatoma was prolonged most after the sequential treatment with D-galactosamine plus 6-azauridine plus 5-fluorouridine. When 6-azauridine was replaced in this combination by N-(phosphonacetyl)-L-aspartate, 80% of ascites hepatoma-bearing rats became tumor-free.

IT 54-25-1

RL: BIOL (Biological study)

Ι

(fluorouridine uptake by neoplasm enhancement by)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 12 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1979:435192 CAPLUS

DN 91:35192

TI Distribution of precursors **labeled** with radioactive isotopes within the early chick embryo

AU Jelinek, R.; Seifertova, M.; Tykva, R.

CS Inst. Exp. Med., Czechoslovak Acad. Sci., Prague, Czech.

SO Colloq. Sci. Fac. Med. Univ. Carol., [Pap.], 21st (1978), 267-71.

Editor(s): Klika, Eduard. Publisher: Univ. Karlova, Prague, Czech.

CODEN: 40MQAS

DT Conference LA English

AB A semiconductog. method was used to det. 3H- and 14C-labeled, precursors along the longitudinal axis of staged, chicken embryos to permit quant, study of nucleic acid and protein synthesis. The method was sensitive enough even to detect specific alterations of DNA, RNA, and protein synthesis following administration of the embryotoxic compds., 6-azauridine and cytosine arabinoside. Thus, thymidine-methyl-3H, uridine-2-14C, leucine-U-3H, and leucine-U-14C were injected into White Leghorn embryos incubated at 37.5.degree., and after 1 h, samples were obtained, mounted on slides and the distribution of the labeled precursors was detd. with a semiconductor detector (fixed surface barrier Si detector) by the method of R. Tykva (1971, 1974). The method was sensitive enough even to detect specific alterations of DNA, RNA, and protein synthesis following administration of the embryotoxic compds., 6-azauridine and cytosine arabinoside.

IT 54-25-1

RL: ANST (Analytical study)
(nucleic acids and proteins formation by chick embryo response to)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 13 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1978:130689 CAPLUS

DN 88:130689

TI Varying distribution of 6-azauridine within the mouse fetoplacental unit

AU Jelinek, R.; Seifertova, M.; Tykva, R.

CS Inst. Exp. Med., Czech. Acad. Sci., Prague, Czech.

SO Folia Morphol. (Prague) (1977), 25(4), 387-95 CODEN: FMORAO; ISSN: 0015-5640

DT Journal

LA English

GΙ

Following i.m. administration of 250 mg (embryotoxic dose) or 25 mg AΒ (nonembryotoxic dose) Riboazauracil (6-azauridine) (I) [54-25-1] together with 100 .mu.Ci I-3H to mice on the 14th day of pregnancy, marked differences in the distribution of radioactivity were obsd. in the embryonic tissue with regard to the high and low doses of I, even though the dose of radiolabeled I was the same. Similar differences in the distribution of labeled I were found when the 25-mg dose was given i.m. and the 250-mg dose was given intraamniotically. embryotoxic dose (250 mg) induced in the placenta accumulation of activity which, 90 min after administration, was over an order of magnitude higher than the sp. activity in the fetus. The activity level in the fetus did not exceed the theor. mean sp. activity in the maternal tissue. The 250-mg dose of I caused extensive malformation in 80% of the fetuses.

54-25-1 TT

RL: PROC (Process) (distribution of, in fetoplacental tissue)

54-25-1 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA CNINDEX NAME)

Absolute stereochemistry.

ANSWER 14 OF 33 CAPLUS COPYRIGHT 2002 ACS 1.6

1977:400161 CAPLUS AN

87:161 DN

Antiviral action and selectivity of 6-azauridine ΤI

Rada, Bretislav; Dragun, Marian ΑU

Inst. Virol., Slovak Acad. Sci., Bratislava, Czech. CS

Ann. N. Y. Acad. Sci. (1977), 284, 410-17 SO

CODEN: ANYAA9 Journal DT

English LΑ

GI

6-Azauridine (I) [54-25-1]-sensitive and -resistant viruses were AB compared with respect to their orotic acid pathways by labeling chick embryo cells with orotic acid-14C during the latent period of viral infection. No differences were detected among vaccinia, Newcastle disease, and vesicular stomatitis viruses. However, I inhibited transport of orotic acid into the cell by 30% and the incorporation of orotic acid into cellular RNA by 50%.

IT 54-25-1

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antiviral activity of)

54-25-1 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA CN INDEX NAME)

Absolute stereochemistry.

ANSWER 15 OF 33 CAPLUS COPYRIGHT 2002 ACS L6

1975:400314 CAPLUS ΑN

DN 83:314

Microsomal inducers of drug-metabolizing enzymes suppress cytidine TInucleotide biosynthesis in rat liver

Seifert, J.; Vacha, J. ΑU

Inst. Pharmacol., Czech. Acad. Sci., Prague, Czech. CS

Arch. Biochem. Biophys. (1975), 167(1), 366-70 SO CODEN: ABBIA4

Journal DT

English LA

For diagram(s), see printed CA Issue. GΙ

Of 17 compds. administered, only substances recognized as inducers of the AB mixed-function oxidases of liver microsomes, such as phenobarbital (I) [50-06-6] decreased the utilization of labeled orotic acid for the synthesis of rRNA cytidine nucleotides in rat liver.

IT 39455-15-7

RL: BIOL (Biological study)

(cytidine nucleotide formation response to)

39455-15-7 CAPLUS RN

ANSWER 16 OF 33 CAPLUS COPYRIGHT 2002 ACS L6

1974:67268 CAPLUS AN

80:67268 DN

Selective action of systemic fungicides and development of resistance TТ

ΑIJ Dekker, J.

Lab. Fytopathol., Wageningen, Neth. CS

Pestic. Chem., Proc. Int. Congr. Pestic. Chem., 2nd (1972), Volume 5, SO 305-8. Editor(s): Tahori, Alexander S. Publisher: Gordon and Breach, New York, N. Y. CODEN: 24WAAY

Conference

LA English

6-Azauridine 5'-phosphate (I) [2018-19-1] (0.06-1.0 .tim.-4M) AB did not inhibit spore germination or germ tube growth in a resistant strain (Type R) of Cladosporium cucumerinum, but orotidine 5'-phosphate [2149-82-8] decarboxylation was completely inhibited. An aq. ext. of ground I-treated spores, however, showed a high orotidine 51-phosphate

decarboxylase [9024-62-8] activity. Intact spores from the resistant strain were less permeable to 14C-labeled 6-azauracil [461-89-2] and 6-azauridine [54-25-1] than spores from the wild (sensitive) strain. Decreased permeability is probably the main factor in the resistance of type R spores to 6-azauracil and its derivs.

IT 2018-19-1

RL: BIOL (Biological study)

(Cladosporium cucumerinum spore germination and growth in response to)

RN 2018-19-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-(5-0-phosphono-.beta.-D-ribofuranosyl)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 54-25-1

RL: BIOL (Biological study)

(Cladosporium cucumerinum spore permeability to)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 17 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1973:500530 CAPLUS

DN 79:100530

TI Effect of 5-azacytidine and 6-azauridine on the synthesis of DNA in embryonic mouse brain mitotic activity and migration of ventricular cells

AU Seifertova, M.; Cihak, A.; Vesely, J.

CS Inst. Org. Chem. Biochem., Czech. Acad. Sci., Prague, Czech.

SO Neoplasma (1973), 20(3), 243-9 CODEN: NEOLA4

DT Journal

LA English

AB 5-Azacytidine (I) [320-67-2] and 6-azauridine (II) [54-25-1] were toxic to the developing fetal brain, as indicated by pycnotic degeneration of ventricular cells and brain malformations in the mouse embryos subsequent to transplacental administration of the agents. II inhibited the uptake of simultaneously administered 3H-labeled thymidine, by the ventricular cells which simultaneously showed decreased mitosis; the pycnotic nuclei lost their capacity to migrate toward the

ventricular surface. In contrast, I-treated fetuses showed ventricular cells heavily labeled with thymidine; the nuclei of these cells were able to migrate toward the ventricular surface and underwent incomplete mitosis followed by pycnotic degeneration. The increased no. of mitotic figures seen after I was due to their abnormal accumulation.

IT 54-25-1

RL: BIOL (Biological study)
 (embryonic brain toxicity of)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 18 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1972:535567 CAPLUS

DN 77:135567

TI Mechanism of resistance of Cladosporium cucumerinum against 6-azauracil and 6-azauridine

AU Dekhuijzen, H. M.; Dekker, J.

CS Inst. Org. Chem., TNO, Utrecht, Neth.

SO Acta Phytopathol. (1971), 6(1-4), 339-43 CODEN: APYBBR

DT Journal

LA English

The wild (N) and resistant (R) strains of C. cucumerinum equally incorporated 14C-labeled 6-azauracil (I) [461-89-2] and showed no difference in the formation of 6-azauridine (II) [54-25-1] from I, but the amt. of 6-azauridine monophosphate (III) [2018-19-1] formed from II was 3-fold lower in the R strain than in the N strain. The poor ability to form III from II by strain R gave rise to a less effective inhibition of orotidine monophosphate decarboxylase [9024-62-8] which consequently resulted in less interference with the incorporation of orotic acid [65-86-1] into RNA. Resistance of strain R to I depended mainly on the failure to form significant amts. of III from II, whereas resistance to III depended on the inability of III to reach the enzyme, orotidine monophosphate decarboxylase, in sufficient amts. Thus, resistance to I and II was attributed to a defect at a stage of uridine kinase [9026-39-5] which normally converts I into II and into III.

IT 54-25-1

RL: PRP (Properties)

(Cladosporium cucumerinum resistance to, mechanism of)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

IT 2018-19-1

RL: FORM (Formation, nonpreparative) (formation of, by Cladosporium cucumerinum in azauracil and azauridine resistance)

RN 2018-19-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-(5-O-phosphono-.beta.-D-ribofuranosyl)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 19 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1972:511017 CAPLUS

DN 77:111017

TI Effect of 6-azauridine in the irradiation of tumors

AU Magdon, E.

CS Inst. Krebsforsch., Berlin, Ger.

SO Radiobiol., Radiother. (1971), 12(4), 535-44 CODEN: RDBGAT

DT Journal

LA German

The effect of 6-azauridine on the radiation-induced retardation of the growth of Yoshida sarcomas, Ehrlich carcinomas, benzopyrene-induced carcinomas, and on the growth and transplantability of spontaneous cancer of the breast was studied. Application of 6-azauridine before irradn. resulted in a significant amplification of the radiation effect. The extent of the amplification was dependent on the interval between 6-azauridine application and irradn. The max. intensifying and sensitizing effect was obtained when 6-azauridine was administered 24 hr before irradn. The radiation sensitizing effect of 6-azauridine is due to a partial synchronization whereby the main part of the cellular population at the time of irradn. is in a phase of max. radiosensitivity. Microautoradiographic examns. show that 24 hr after the application of 6-azauridine the incorporation of thymidine-3H into the DNA of ascites cells of Ehrlich-carcinoma is increased compared with that for the untreated cells. Since the labeling index is increased by the application of 6-azauridine, the early S-phase is the cellular stage that is responsible for the radiation sensitizing effect after a blockage of the G1 phase.

IT 54-25-1

RL: BIOL (Biological study)
(neoplasm response to radiation in relation to)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 20 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1972:82108 CAPLUS

DN 76:82108

TI Induction of tryptophan oxygenase in Cicer arietinum seedlings by indole-3-acetic acid and cyclic 3',5'-adenosine monophosphate

AU Azhar, S.; Krishna Murti, C. R.

CS Div. Biochem., Cent. Drug Res. Inst., Lucknow, India

SO Indian J. Biochem. Biophys. (1971), 8(4), 210-13 CODEN: IJBCAS

DT Journal

LA English

AΒ Tryptophan oxygenase activity was not detected in dormant seeds of Bengal gram, Cicer arietinum, but appeared after 48 hr of germination, reached a peak value in 120 hr, and remained unchanged thereafter. Seedlings aged 48, 72, or 96 hr exhibited a 2- to 3-fold increase in enzyme activity on preincubation in a medium contg. 1 .tim. 10-6M IAA [87-51-4] or 1 .tim. 10-4M cyclic AMP [60-92-4]. Beyond 96 hr of germination, IAA or cyclic AMP did not exert this stimulation of enzyme activity in vitro. IAA-mediated stimulation of activity in 48 hr or 72 hr seedlings was inhibited 50% by 1 .tim. 10-3M cycloheximide (I) [66-81-9], 1 .tim. 10-2M DL-ethionine [67-21-0], 1 .tim. 10-2M azauridine [54-25-1], or 1 .tim. 10-3M DL-p-fluorophenylalanine (II) [51-65-0]. Incorporation of 14C-labeled valine [72-18-4] into TCA precipitable proteins of 72 hr seedlings was stimulated by 1 .tim. 10-6M IAA. The in vitro enzyme stimulation by 1 .tim. 10-4M cyclic AMP in 72 hr seedlings was also inhibited by 1 .tim. 10-2M I or 1 .tim. 10-3M II.

IT 54-25-1

RL: BIOL (Biological study)

(indoleacetic acid induced tryptophan oxygenase activity in chick pea inhibition by)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CF INDEX NAME)

L6 ANSWER 21 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1972:21183 CAPLUS

DN 76:21183

TI Transfer of 6-azauridine through the placental barrier in the rat

AU Gutova, M.; Elis, J.; Raskova, H.

CS Fac. Pediatr., Charles Univ., Prague, Czech.

SO Neoplasma (1971), 18(5), 529-31 CODEN: NEOLA4

DT Journal

LA English

AB 6-Azauridine (I) [54-25-1] was readily and efficiently penetrated the placental barrier in rats on day 16 and 20 of gestation, supporting the high teratogenic and embryotoxic effects. Peak levels in the placenta and fetuses were obsd. 40-80 min after the i.v. injection of 4,5-14C-labeled I. The radioactivity in fetuses accounted for 35-36% of that in maternal plasma. The rate of penetration was about the same at both stages of gestation.

IT 54-25-1

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study) (placental transport and teratogenicity of)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 22 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1972:130 CAPLUS

DN 76:130

TI Pharmacokinetics of 6-azauridine-4,5-14C in rats with adjuvant-induced polyarthritis

AU Perlik, F.; Elis, J.

CS Inst. Pharmacol., Czech. Acad. Sci., Prague, Czech.

SO Physiol. Bohemoslov. (1971), 20(2), 181-4 CODEN: PHBOAP

DT Journal

LA English

AB The blood, brain, kidney, and small intestine levels of 14C-labeled 6-azauridine (I) [54-25-1] were similar in control rats and in those with adjuvant-induced arthritis. Higher levels of antimetabolite were detected in the spleen of adjuvant rats for 15 min and liver for the full 90-min exptl. period. The rate of disappearance of I from the livers of control and adjuvant rats was almost the same. There may be more binding of I in the liver of adjuvant rats, but the reasons for this are not yet clear.

IT 54-25-1

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (metabolism of, in arthritis)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 23 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1971:401782 CAPLUS

DN 75:1782

TI Role of pharmacologically active nucleoside derivatives in RNA translation

AU Skoda, Jan

CS Inst. Org. Chem. Biochem., Czech. Acad. Sci., Prague, Czech.

SO Biochem. Aspects Antimetab. Drug Hydrozylation, Fed. Eur. Biochem. Soc., Meet., 5th (1969), Meeting Date 1968, 23-30. Editor(s): Shugar, D. Publisher: Academic, London, Engl.

CODEN: 22ZEAD

DT Conference

LA English

With a uridylic acid (I) and 6-azacytidylic acid (II) ratio of 3:1, slight amts. of phenylalanine-14C were incorporated into Escherichia coli in vitro. These copolymers had no messenger activity for any amino acid which included I or cytidylic acid (III) in its codon. II could not replace III or I in the codon. In the presence of I and II, the anomalous polyribonucleotide significantly enhanced the messenger activity of poly U for phenylalanine. The replacement of any pyrimidine ribonucleoside in the valine codon by 6-azauridine (IV) or 6-azacytidine (V) resulted in a nonfunctional unit. Binding of 14C-labeled valyl-tRNA to ribosomes was neg. even with triplets contg. IV or V in the 3rd position corresponding to that of inosine in the valine anticodon. Incorporation of 6-azapyrimidines into nucleic acids probably did not cause errors in the transfer of genetic information. Triplets contg. arabinose residues did not stimulate binding.

IT 32972-70-6

RL: BIOL (Biological study)

(valyl-transfer ribonucleic acid binding to ribosomes in response to)

RN 32972-70-6 CAPLUS

CN Guanosine, uridylyl-(5'.fwdarw.3')-6-azauridylyl-(5'.fwdarw.3')- (8CI) (CA INDEX NAME)

IT 32972-69-3

RL: BIOL (Biological study)

(valy1-transfer ribonucleic acid binding to ribosomes stimulation by)

RN 32972-69-3 CAPLUS

CN Guanosine, 6-azauridylyl-(5'.fwdarw.3')-uridylyl-(5'.fwdarw.3')- (8CI) (CA INDEX NAME)

- L6 ANSWER 24 OF 33 CAPLUS COPYRIGHT 2002 ACS
- AN 1971:96895 CAPLUS
- DN 74:96895
- TI Effect of 6-azaridine on the labeling index and the mean grain count of Ehrlich-carcinoma ascites cells after [3H]-thymidine incubation AU Magdon, Erwin; Delzer, Wilfried

Inst. Krebsforsch., Dsch. Akad. Wiss. Berlin, Berlin-Buch, Ger. CS

Arch. Geschwulstforsch. (1970), 36(3), 247-52 SO CODEN: ARGEAR

Journal DТ

LA German

The incorporation of thymidine-3H into Ehrlich ascitic carcinoma cells in AB mice 24 hr after i.p. injection of 6-azauridine was examd. At this time the labeling index of the carcinoma cells was increased 1.5-fold. The result sindicated that this increase was due to more cells entering the early S phase of the cell cycle and that the accumulation of cells in this phase is responsible for the radiosensitizing effect of 6-azauridine.

TΤ 54-25-1

> RL: BIOL (Biological study) (thymidine metabolism by carcinoma in response to)

RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA CN INDEX NAME)

Absolute stereochemistry.

ANSWER 25 OF 33 CAPLUS COPYRIGHT 2002 ACS L6

AN1970:531271 CAPLUS

DN 73:131271

Preparation of nucleotides and oligonucleotides by thermal reaction TI

Moravek, Josef; Skoda, Jan IN

SO Czech., 5 pp.

CODEN: CZXXA9

DT Patent

Czech LΑ

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE ____ ----------19691015 CS CS 133464 19651202

PΙ AB Labeled title compds. are obtained by thermal phosphorylation of nucleosides which are heated at 155-65.degree. with an alk. metal phosphate or nucleoside 2'(3')-phosphate in the ratio 1:5. The mixt. is chromatographed and the products isolated from the eluate. Treating labeled nucleosides (14C, 3H, 35S) with nonradioactive phosphates gives compds. contg. soft .beta.-emitters in the nucleoside moiety, reaction of unlabeled nucleosides with 32P-phosphates gives compds. contg. a hard .beta.-emitter in the phosphate residue. The following nucleosides were phosphorylated: cytidine, uridine, pseudouridine, orotidine, deoxycytidine, deoxyuridine, thymidine; (with a change in the base) 6-azauridine, 6-azacytidine; (with a substitution on the base) 5-hydroxyuridine; (with a change in the sugar) N-xylosyluracil; (antibiotics) tubercidine.

IT 29838-80-0

RL: RCT (Reactant)

(reaction of, with labeled sodium phosphate)

RN29838-80-0 CAPLUS

CN as-Triazine-5,6-14C2-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (8CI) Absolute stereochemistry.

L6 ANSWER 26 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1970:118702 CAPLUS

DN 72:118702

6-Azauridine as an inhibitor of the synthesis of Herpesvirus hominis ΤI

ΑU Falke, Dietrich; Rada, B.

CS Inst. Med. Microbiol., Johannes Gutenberg Univ., Mainz, Ger.

SO Acta Virol. (Prague), Engl. Ed. (1970), 14(2), 115-23

CODEN: AVIRA2

DTJournal

LA English

6-Azauridine (AzUR) (10 mg/ml) added immediately after the infection of a AB rabbit kidney cell system with giant cell-forming Herpesvirus strains, inhibited the virus yield at 20 hr postinfection; this action of AzUR was more pronounced when cells were starved before infection. The addn. of uridine or cytidine completely reversed the AzUR inhibition, whereas thymidine or orotic acid did not alter the inhibitory effect of this compd., with respect to both giant cell formation and synthesis of infective particles. AzUR completely inhibited the Herpesvirus-induced giant cell formation when added until 90 min postinfection; later addn. of this antimetabolite exerted only a minor effect. Actinomycin C (1.5 .mu.g/ml) blocked the appearance of giant cells when add ed until 2 hr postinfection. Thus, RNA (probably m-RNA) synthesis is necessary before giant cell formation can be initiated. To correlate the early inhibitor-sensitive step indicated by giant cell fo rmation to the synthesis of infective particles, infected cultures were tr eated with cytosine arabinoside (100 .mu.g/ml) or cycloheximide (1.25 .mu.g/ml) at various times after infection. Drugs added up to .apprx.5 hr postinfection inhibited synthesis of infective virus particles. AzUR added to cultures at 150 min postinfection had no effect on giant cell formation; however, the addn. of this compd. plus cycloheximide completely inhibited the appearance of giant cells. AZUR reduced the incorporation of tritium-labeled thymidine into DNA of uninfected and virus-infected cells by 66% and 82%, resp. The redn. in viral DNA synthesis was attributed to the synthesis of new m-RNA before the beginning of viral DNA synthesis. Similar results were obtained with azaquanine. 54-25-1

ΙT

RL: PROC (Process)

(virucidal action of, to herpes simplex virus)

RN54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) INDEX NAME)

L6 ANSWER 27 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1970:108077 CAPLUS

DN 72:108077

TI Role of uridine in the synthesis of phage-specific DNA in the cells of Escherichia coli infected with phage T2

AU Pravdina, N. F.; Galegov, G. A.

CS Inst. Virol., Moscow, USSR

SO Biokhimiya (1970), 35(1), 85-8 CODEN: BIOHAO

DT Journal

LA Russian

The role of 14C-labeled uridine in the synthesis of nucleic acids was studied in E. coli B cells infected with phage T2 to clarify the role of uridine kinase in the development of phage infection.

Incorporation of the nucleosides into RNA decreased sharply during phage infection. DNA of infected bacteria showed a 3-fold increase in labeled uridine compared with the control. Preliminary 90-min pre-incubation of bacteria with 1000 .mu.g/ml 6-azauridine decreased incorporation of 14C-labeled uridine into RNA in noninfected bac-teria, but the ability to incorporate uridine into RNA in the in-fected cells was close to the same level in the presence and ab-sence of antimetabolite. 6-Azauridine inhibited incorporation of labeled uridine and thymidine into phage-induced DNA.

IT 54-25-1

RL: BIOL (Biological study)

(deoxyribonucleic acid formation by bacteriophage-infected Escherichia coli in presence of)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 28 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1969:419396 CAPLUS

DN 71:19396

TI Biochemically important **labeled** compounds. V. Biosynthetic preparation of 6-azauridine-4,5-14C and its isolation by gel filtration

AU Moravek, Josef; Skoda, Jan

CS Ceskoslov. Akad. Ved, Prague, Czech.

SO Collect. Czech. Chem. Commun. (1969), 34(6), 1837-40 CODEN: CCCCAK

DT Journal

LA English

AB 6-Azauracil-4,5-14C (I) is added to a medium contg. Escherichia coli (CA 51: 13056b) in the logarithmic phase of growth to obtain a 2 .times. 10-3M I. The fermentation is continued another 14 hrs. at 37.degree., the medium chilled, the bacterial mass and pptd. orotic acid centrifuged, the supernatant concd. in vacuo and filtered. The title compd. is sepd. in a 70% yield on a column of Biogel P-2 (200-400 mesh) in distd. water from unchanged I which is used in the next fermentation batch. The overall conversion is practically quant.

IT 25541-14-4

RL: FORM (Formation, nonpreparative) (formation of, by Escherichia coli)

RN 25541-14-4 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione-5,6-14C2, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 29 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1968:400055 CAPLUS

DN 69:55

TI Effect of 5'-terminal phosphate on the recognition of some dinucleoside phosphates by aminoacyl-14C-tRNA

AU Gruenberger, Dezider; Holy, Antonin; Sorm, Frantisek

CS Ceskoslov. Akad. Ved, Prague, Czech.

SO Biochim. Biophys. Acta (1968), 157(2), 439-42 CODEN: BBACAQ

DT Journal

LA English

AΒ

5'-Terminal phosphate increased not only the template activity of the triplet codon for valine, but also that of the dinucleoside phosphate. Whereas GpU was without effect, pGpU stimulated the binding of 14Clabeled valy1-tRNA to ribosomes. Similarly, pGpUpU had a markedly higher stimulatory effect than GpUpU. On the other hand, 3'-terminal phosphate, pGpUp, decreased the template activity of the dinucleotide. Dinucleotides with 5'-phosphate and contg. 5-bromomethyluridine BrU or 5-methyluridine instead of uridine stimulated the binding of valyl-tRNA to ribosomes. The increased template activity of pGpBrU in comparison with pGpU could be anticipated on the basis of greater stability of the binding between halogen derivs. of uridine and the complementary nucleosides in the anticodon part of the tRNA mol. In contrast, by substitution of a Me group in the N-3-position of uridine N-3-MeU for the H atom involved in the codon-anticodon base pairing, the pGpN-3-MeU formed was totally inactive in this system. Similarly, substitution of 6-azauridine for uridine abolished the template activity of the nucleotide. However, the 5'-terminal phosphate did not enhance the effect of the appropriate doublet or triplet on the binding of aspartyl-tRNA or glutamyl-tRNA to ribosomes. These results demonstrated that a dinucleotide with 5'-terminal phosphate may only occasionally be recognized by

aminoacyl-tRNA; this is possibly related to the evolution of the code as was proposed by Rottman and Nirenberg (1966).

IT 20512-61-2

> RL: BIOL (Biological study) (as template for valyl-ribonucleic acid)

20512-61-2 CAPLUS RN

6-Azauridine, 5'-O-phosphonoguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX CN

Absolute stereochemistry.

ANSWER 30 OF 33 CAPLUS COPYRIGHT 2002 ACS L6

1968:36337 CAPLUS ΑN

68:36337 DN

Thyrotropin stimulation of pyrimidine nucleotide Synthesis in bovine TТ thyroid

Lindsay, Raymond H.; Cash, Anna G.; Hill, Johnie Banks ΑU

Univ. of Alabama Med. Center, Birmingham, Ala., USA CS

Biochem. Biophys. Res. Commun. (1967), 29(6), 850-5 SO CODEN: BBRCA9

DТ Journal

LA English

One and 20 micromoles of azauridine per 3 cc. vol. inhibited by 85.1 and AB 98.4%, resp., the formation of 14CO2 from orotic acid-carboxyl-14C in bovine thyroid slices. However, 20 micromoles of azauridine had no effect on the oxidn. of glucose-14C. Thyrotropin (1 unit 3 cc.) enhanced by 30% the conversion of the labeled orotic acid to pyrimidine nucleotide in the absence of azauridine but had no effect in the presence of 5 micromoles azauridine. The thyrotropin stimulation occurred both in the presence and in the absence of glucose (4 micromoles). Labeled CO2 production from orotic acid-carboxyl-14C in bovine thyroid slices is due almost exclusively to the formation of pyrimidine nucleotides. Thyrotropin stimulates total pyrimidine nucleotide synthesis in bovine thyroid slices. These findings along with previously reported findings strongly indicate that 1 site at which thyrotropin may affect nucleic acid synthesis is the formation of nucleotides.

IT 54-25-1

RL: BIOL (Biological study) (pyrimidine nucleotide formation inhibition by, in thyroid gland, thyrotropin in relation to)

54-25-1 CAPLUS RN

1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA CN INDEX NAME)

L6 ANSWER 31 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1967:508969 CAPLUS

DN 67:108969

TI Nucleoside 5'-polyphosphates and .alpha.,.omega.-bis-(nucleoside-5')polyphosphates

IN Moffatt, John G.

PA Syntex Corp.

SO U.S., 10 pp.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PΙ 19670323 US 19650317 US 3321463 A process is claimed for the prepn. of the title compds. which comprises treating, under anhyd. conditions in a dialkyl sulfoxide, a nucleoside 5'-polyphosphate terminal phosphoramidate tertiary amine salt with a tertiary amine salt of an inorg. phosphate or a nucleoside 5'-phosphate. Little tendency for the polyphosphates to degrade to lower phosphates is encountered in this process in contrast to the general method using pyridine solvent. Na4P2O7.10H2O (1 millimole) in 10 ml. H2O was passed through a column of Dowex 50W-X8 ion exchange resin (C5H5N form). The effluent and a 30-ml. H2O wash was evapd. in vacuo to a 10 ml. vol. C5H5N (30 ml.) and 4.2 millimoles Bu3N were added and the homogeneous soln. was evapd. to a sirup and dried by four successive evapns. in vacuo with 10-ml. portions dry C5H5N and two evapns. with 5-ml. portions dry C6H6. The pyrophosphate was dissolved in 4 sep. 1 ml. portions dry Me2SO and added successively to 0.25 millimole dry 4-morpholino-N,N'dicyclohexylcarboxamidinium salt of 2'-deoxyadenosine 5'phosphoromorpholidate. The resulting clear soln. was kept 4 days at room temp. (the disappearance of the phosphoromorpholidate was followed by paper chromatog.). The mixt. was treated with 30 ml. H2O and the soln. chromatographed on DEAE-cellulose (HCO3-form). The pooled triphosphate fraction was evapd. to dryness at 30-5.degree. and (Et3NH)HCO3 removed by evapn. with MeOH. The residue was dissolved in 5 ml. MeOH and treated with a N soln. (6 equivs.) of NaI in Me2CO and 75 ml. Me2CO. The ppt. was collected by centrifugation, washed with Me2CO and dried over P2O5 to give the Na salt of 2'-deoxyadenosine 5'-triphosphate as a white chromatographically homogeneous powder. Similarly prepd. in 75-80% yields were the Na salts of 5'-triphosphate: 2'-deoxyguanosine, 6-azauridine, cytidine, 2'-deoxycytidine, and thymidine. Dicyclohexylcarbodiimide (5 millimoles) in 35 ml. tert-BuOH was added over 9 hrs. to a refluxing soln. of 1 millimole of the morpholine salt of ADP and 2.4 millimoles morpholine in 20 ml. aq. 50% tert-BuOH. Refluxing was continued until paper chromatog. showed the absence of ADP. The mixt. was cooled and filtered, and the filtrate evapd. in vacuo. The aq. conc. was washed by extn. with Et2O and, after adjusting to pH 8, was chromatographed to give 83%P1-(adenosine-5')-P2-(4'-morpholino)diphosphate as the Et3NH salt (I). Substituting NH4OH for morpholine gave the Et3NH salt of P1-(adenosine-5')-P2-aminotriphosphate. I was dissolved in a MeOH soln.

of the 4-morpholine salt of N,N'-dicyclohexylcarbodiimide. The soln. was evapd. to dryness and the residue dissolved in MeOH and pptd. with Et2O to give the 4-morpholine N,N'-dicyclohexylcarboxamidinium salt (II) of I. Similarly prepd. were the Et3NH salts of P1-(adenosine-5')-P3-(4'morpholino)triphosphate (III) and P1-(uridine-5')-P3-(4'morpholino)phosphate (IV) and the 4-morpholino-N,N'dicyclohexylcarboxamidinium salts (V) of III and IV. I was mixed with an excess of M soln. of CaCl2 in EtOH to ppt. the Ca salt of I. II (0.25 millimole) and 1 millimole of Bu3NH orthophosphate (VI) in 3 ml. Me2SO was heated 24 hrs. at 40.degree.. The mixt. was dild. with 40 ml. H2O and chromatographed on DEAE-cellulose (HCO3-form) to give 71% ATP (isolated as the Na salt). II (0.1 millimole) and 0.3 millimole of 32P-labeled VI (contg. 2 .mu.c. of 32P) in 2 ml. Me2SO was heated 45 hrs. at 35.degree. and worked up to yield the Na salt of ATP-.gamma.-32P. millimole) and 0.4 millimole of the Bu3NH salt of AMP in 2 ml. Me2SO was kept 4 days at room temp. to give 31% of the Na salt of .alpha., .zeta.-bis(adenosine-5')tetraphosphate.

IT 18423-44-4P

RN 18423-44-4 CAPLUS

CN as-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl-, 5'-(tetrahydrogen triphosphate), sodium salt (8CI) (CA INDEX NAME)

Absolute stereochemistry.

●x Na

L6 ANSWER 32 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1967:113266 CAPLUS

DN 66:113266

TI Action of azauracil derivatives on Saccharomyces cerevisiae. II.

Differences in metabolic behavior between normal and 6-azauracil-resistant
strains

AU Wiesenfeld, Mireille; Crokaert, Robert

CS Fac. Med. Univ. Libre, Brussels, Belg.

SO Bull. Soc. Chim. Biol. (1967), 49(2), 191-203 CODEN: BSCIA3

DT Journal

LA French

AB cf. CA 66, 53169f. The uptake of 6-azauracil-2-14C (I) by normal S. cerevisiae cells rapidly reached a max. by the 60th-120th min. of incubation, then slowly declined. The uptake by a 6-azauracil-resistant mutant rapidly reached a plateau, which corresponded to 11% of the max. uptake observed in the normal strain. Similarly, the uptake of uracil-2-14C (II) by the resistant strain was only 0.2% that of the normal strain, although the uptake increased with incubation time in both strains. Unlabeled compds. chromatographically similar to orotic acid (or orotidine) and labeled 6-azauridine appeared in the medium after incubation of the normal strain with I, but not after incubation of the

resistant strain. One and 2 unidentified labeled compds. were detected in the medium after incubation of the resistant and normal strains, resp., with II. Acid-sol. exts. of normal cells grown on I contained I, 6-azauridine, and an unidentified compd., while exts. from the resistant strain contained only I. Acid-sol. exts. of normal cells contained II and 3 unidentified labeled compds. after incubation with II, while the resistant strain contained only II. Uracil (0.5 micromole/ml.), dihydrouracil (1.0 micromole/ml.), and N-carbamoyl-.beta.-alanine (2.5 micromoles/ml.) reduced the uptake of I by the normal cells by 85, 64, and 35%, resp. II was incorporated into the RNAs of the normal but not the resistant strain. The specific activity of uridine phosphorylase was similar in both strains, excluding a loss of this enzyme in the mutant. 15 references.

IT 54-25-1

RL: BIOL (Biological study)

(as 6-azauracil metabolite in Saccharomyces cerevisiae)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 33 OF 33 CAPLUS COPYRIGHT 2002 ACS

AN 1967:45237 CAPLUS

DN 66:45237

TI Relation between the metabolic effects and the pregnancy-interrupting property of 6-azauridine in mice

AU Raska, Karel, Jr.; Zedeck, Morris S.; Welch, Arnold D.

CS Sch. of Med., Yale Univ., New Haven, Conn., USA

SO Biochem. Pharmacol. (1966), 15(12), 2136-8 CODEN: BCPCA6

DT Journal

LA English

When administered soon after implantation of the fertilized ovum, a single AB i.p. injection of 6-azauridine (I) (500 mg./kg.) into mice resulted in complete resorption of the embryo, whereas, when administered during the 2nd half of pregnancy, even repeated injection of I did not consistently produce an interruption of pregnancy. To det. the difference between the effects of I upon the early and late stages of fetal development, the metabolic transformation and biochem. effects of I were compared in 6-day embryos and in 15-day fetuses of mice. Injections of I inhibited the incorporation of labeled orotic acid into nucleic acids to the same extent at both days 6 and 15 of fetal development. In vitro, I (5 .times. 10-5M) inhibited orotidylate decarboxylase activity in particle-free supernatant fractions to the same extent in both the fetus and embryo. The time difference in the pregnancy-interrupting effect of I thus reflects a difference in sensitivity of embryonic and fetal tissues to inhibition of nucleic acid synthesis.

IT 54-25-1

RL: BIOL (Biological study)

(abortion from, nucleic acid formation and)

RN 54-25-1 CAPLUS

CN 1,2,4-Triazine-3,5(2H,4H)-dione, 2-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

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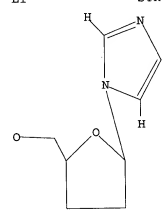
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FILE COVERS 1907 - 25 Sep 2002 VOL 137 ISS 13 FILE LAST UPDATED: 23 Sep 2002 (20020923/ED)

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=> s 12

L3 85 L2

=> s 13 and label

48825 LABEL

L4 0 L3 AND LABEL

=> s 13 and label?

374492 LABEL?

L5 15 L3 AND LABEL?

=> d l15 bib abs hit str 1-15

L15 NOT FOUND

The L-number entered has not been defined in this session, or it has been deleted. To see the L-numbers currently defined in this session, enter DISPLAY HISTORY at an arrow prompt (=>).

=> d 15 bib abs hitstr 1-15

L5 ANSWER 1 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 2001:851808 CAPLUS

DN 135:367666

TI Nucleotide analogs and their use in **labeling** nucleic acids for hybridization assays

IN McGall, Glenn; Barone, Anthony D.

PA USA

SO U.S. Pat. Appl. Publ., 47 pp., Cont.-in-part of U.S. Appl. 2001 18,514. CODEN: USXXCO

DT Patent LA English

FAN.CNT 2

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|----------------|------|----------|-----------------|----------|
| | | | | | |
| ΡI | US 2001044531 | A1 | 20011122 | US 2001-780574 | 20010209 |
| | US 2001018514 | A1 | 20010830 | US 1998-126645 | 19980731 |
| PRAI | US 1998-126645 | A2 | 19980731 | | |

OS MARPAT 135:367666

AB Nucleic acid labeling compds. contg. heterocyclic derivs. are disclosed. The heterocyclic deriv. contg. compds. are synthesized by condensing a heterocyclic deriv. with a cyclic group (e.g. a ribofuranose deriv.). The labeling compds. are suitable for enzymic attachment to a nucleic acid, either terminally or internally, to provide a mechanism of nucleic acid detection. Thus, a no. of biotin- or fluorescein purine- and pyrimidine-.beta.-D-ribofuranoside analogs were prepd. These analogs were successfully incorporated into hybridization probes (using terminal deoxynucleotidyltransferase) and utilized in single nucleotide polymorphism geno-typing using micro-chip arrays.

IT 257297-78-2P 257297-98-6P 373390-75-1P 373391-24-3P 373391-42-5P 373391-43-6P

RL: BSU (Biological study, unclassified); IMF (Industrial manufacture); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (nucleotide analogs and their use in labeling nucleic acids for hybridization assays)

RN 257297-78-2 CAPLUS

CN Triphosphoric acid, P-[[(2S,5R)-5-[4-[[[4-[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]-1-oxohexyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2-furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RN 257297-98-6 CAPLUS

Triphosphoric acid, P-[[(2S,5R)-5-[4-[[[4-[[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5yl)carbonyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2furanyl]methyl] ester (9CI) (CA INDEX NAME)

OPO₃H₂
OPO₃H₂
O
N
HO
O
$$(CH_2)_4$$
H
O
O

PAGE 1-B

373390-75-1 CAPLUS RN

Spiro[isobenzofuran-1(3H),9'-[9H]xanthene]-5-carboxamide,
3',6'-dihydroxy-3-oxo-N-[4-[[[1-[(2R,5S)-tetrahydro-5-(hydroxymethyl)-2-furanyl]-1H-imidazol-4-yl]carbonyl]amino]butyl]- (9CI) (CA INDEX NAME) CN

RN 373391-24-3 CAPLUS

CN Spiro[isobenzofuran-1(3H),9'-[9H]xanthene]-5-carboxamide, 3',6'-dihydroxy-3-oxo-N-[4-[[[1-[(2S,5S)-tetrahydro-5-(hydroxymethyl)-2-furanyl]-1H-imidazol-4-yl]carbonyl]amino]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

__OH

RN 373391-42-5 CAPLUS

CN Triphosphoric acid, P-[[(2S,5S)-5-[4-[[[4-[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]-1-oxohexyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2-furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RN 373391-43-6 CAPLUS

CN Triphosphoric acid, P-[[(2S,5S)-5-[4-[[[4-[[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5-

yl)carbonyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A HO_

PAGE 1-B

257297-73-7P 257297-74-8P 257297-75-9P IT

257297-76-0P 257297-77-1P

RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (nucleotide analogs and their use in labeling nucleic acids for hybridization assays)

257297-73-7 CAPLUS RN

1H-Imidazole-4-carboxylic acid, 1-[(2R,5S)-5-[[[(1,1-CNdimethylethyl)dimethylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

257297-74-8 CAPLUS RN

CN 1H-Imidazole-4-carboxamide, N-(4-aminobutyl)-1-[(2R,5S)-5-[[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]tetrahydro-2-furanyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 257297-75-9 CAPLUS

CN 1H-Imidazole-4-carboxamide, 1-[(2R,5S)-5-[[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]tetrahydro-2-furanyl]-N-[4-[(trifluoroacetyl)amino]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$F_3C \xrightarrow{N}_H (CH_2)_4 \xrightarrow{N}_H N \xrightarrow{N}_R S \xrightarrow{N}_S i$$
 Bu-t

RN 257297-76-0 CAPLUS

CN 1H-Imidazole-4-carboxamide, 1-[(2R,5S)-tetrahydro-5-(hydroxymethyl)-2-furanyl]-N-[4-[(trifluoroacetyl)amino]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 257297-77-1 CAPLUS

CN Triphosphoric acid, P-[[(2S,5R)-tetrahydro-5-[4-[[[4-[(trifluoroacetyl)amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]-2-furanyl]methyl] ester (9CI) (CA INDEX NAME)

$$F_{3}C$$

$$N$$

$$H$$

$$N$$

$$N$$

$$R$$

$$S$$

$$O$$

$$OH$$

$$OPO_{3}H_{2}$$

$$O$$

$$OH$$

$$OPO_{3}H_{2}$$

$$OH$$

$$OH$$

L5 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 2001:675161 CAPLUS

DN 136:37868

TI Novel nucleoside triphosphate analogs for the enzymatic labeling of nucleic acids

AU Barone, A. D.; Chen, C.; McGall, G. H.; Rafii, K.; Buzby, Philip R.; Dimeo, James J.

CS Affymetrix, Inc., Santa Clara, CA, USA

SO Nucleosides, Nucleotides & Nucleic Acids (2001), 20(4-7), 1141-1145 CODEN: NNNAFY; ISSN: 1525-7770

PB Marcel Dekker, Inc.

DT Journal

LA English

We have evaluated several novel nucleotide analogs suitable for enzymic labeling of nucleic acid targets for a variety of array-based assays. Two new reagents in particular, a C4-labeled 1-(2',3'-dideoxy-.beta.-D-ribofuranosyl) imidazole-4-carboxamide 5'-triphosphate and an N1-labeled 5-(.beta.-D-ribofuranosyl)-2,4(1H,3H)-pyrimidinedione 5'-triphosphate, were found to be excellent substrates for labeling with terminal deoxynucleotidyl transferase and T7 RNA polymerase, resp.

IT 257297-98-6 380601-34-3

RL: BCP (Biochemical process); BIOL (Biological study); PROC (Process) (prepn. of nucleoside triphosphate analogs for enzymic labeling of nucleic acids)

RN 257297-98-6 CAPLUS

CN Triphosphoric acid, P-[[(2S,5R)-5-[4-[[[4-[[(3',6'-dihydroxy-3oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5yl)carbonyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

HO_

RN 380601-34-3 CAPLUS

CN Triphosphoric acid, P-[[(2S,5R)-5-[4-[[[4-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2-furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L5 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2002 ACS
- AN 2000:98825 CAPLUS
- DN 132:133201
- TI Nucleotide analogs and their use in **labeling** nucleic acids for hybridization assays
- IN McGall, Glenn H.; Barone, Anthony D.
- PA Affymetrix, Inc., USA
- SO PCT Int. Appl., 69 pp. CODEN: PIXXD2
- DT Patent

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English
LA
FAN.CNT 2
                                                            DATE
                                           APPLICATION NO.
                      KIND DATE
     PATENT NO.
                                           _____
                      _ _ _ _
                            _ _ _ _ _ _ _ _
                                           WO 1999-US12390 19990720
                      A2
                            20000210
PΙ
     WO 2000006771
        W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP,
             KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,
             MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM,
             TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
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             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                           US 1998-126645
                                                             19980731
     US 2001018514
                       A1
                            20010830
                                                             19990720
                                           AU 1999-52035
     AU 9952035
                       Α1
                            20000221
                                           EP 1999-937150
                                                             19990720
                       A2
                            20010822
     EP 1124838
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                                           JP 2000-562553
                                                             19990720
                       T2
                            20020716
     JP 2002521495
PRAI US 1998-126645
                       Α
                            19980731
     WO 1999-US12390
                       W
                            19990720
OS
     MARPAT 132:133201
     Nucleic acid labeling compds. contg. heterocyclic derivs. are
AB
     disclosed. The heterocyclic deriv. contg. compds. are synthesized by
     condensing a heterocyclic deriv. with a cyclic group (e.g. a ribofuranose
     deriv.). The labeling compds. are suitable for enzymic
     attachment to a nucleic acid, either terminally or internally, to provide
     a mechanism of nucleic acid detection. Thus, a no. of biotin- or
     fluorescein purine- and pyrimidine-.beta.-D-ribofuranoside analogs were
            These analogs were successfully incorporated into hybridization
     probes (using terminal deoxynucleotidyltransferase) and utilized in single
     nucleotide polymorphism genotyping using microchip arrays.
     257297-78-2P 257297-98-6P
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); SPN
     (Synthetic preparation); BIOL (Biological study); PREP (Preparation); PROC
     (Process)
        (nucleotide analogs and their use in labeling nucleic acids
        for hybridization assays)
     257297-78-2 CAPLUS
RN
     Triphosphoric acid, P-[[(2S,5R)-5-[4-[[[4-[[6-[[5-[(3aS,4S,6aR)-hexahydro-
CN
     2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]-1-
```

Absolute stereochemistry.

PAGE 1-A

oxohexyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2-

furanyl]methyl] ester (9CI) (CA INDEX NAME)

RN 257297-98-6 CAPLUS

CN Triphosphoric acid, P-[[(2S,5R)-5-[4-[[[4-[[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5-yl)carbonyl]amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]tetrahydro-2-furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A HO

PAGE 1-B

IT 257297-73-7P 257297-74-8P 257297-75-9P

257297-76-0P 257297-77-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(nucleotide analogs and their use in **labeling** nucleic acids for hybridization assays)

RN 257297-73-7 CAPLUS

CN 1H-Imidazole-4-carboxylic acid, 1-[(2R,5S)-5-[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 257297-74-8 CAPLUS

CN 1H-Imidazole-4-carboxamide, N-(4-aminobutyl)-1-[(2R,5S)-5-[[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]tetrahydro-2-furanyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$H_2N$$
 $(CH_2)_4$
 N
 N
 R
 S
 O
 $Bu-t$

RN 257297-75-9 CAPLUS

CN 1H-Imidazole-4-carboxamide, 1-[(2R,5S)-5-[[[(1,1-dimethylethyl)dimethylsilyl]oxy]methyl]tetrahydro-2-furanyl]-N-[4-[(trifluoroacetyl)amino]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 257297-76-0 CAPLUS

CN 1H-Imidazole-4-carboxamide, 1-[(2R,5S)-tetrahydro-5-(hydroxymethyl)-2-furanyl]-N-[4-[(trifluoroacetyl)amino]butyl]- (9CI) (CA INDEX NAME)

RN 257297-77-1 CAPLUS

CN Triphosphoric acid, P-[((2S,5R)-tetrahydro-5-[4-[[[4-[(trifluoroacetyl)amino]butyl]amino]carbonyl]-1H-imidazol-1-yl]-2furanyl]methyl] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1995:715348 CAPLUS

DN 123:103310

TI Imidazoleacetic acid, a .gamma.-aminobutyric acid receptor agonist, can be formed in rat brain by oxidation of histamine

AU Thomas, Boban; Prell, George D.

CS Dep. Pharmacol., Mt. Sinai Sch. Med. City Univ. New York, New York, NY, USA

SO Journal of Neurochemistry (1995), 65(2), 818-26 CODEN: JONRA9; ISSN: 0022-3042

PB Lippincott-Raven

DT Journal

LA English

It is generally accepted that in mammalian brain histamine is metabolized AB solely by histamine methyltransferase (HMT), to form tele-methylhistamine, then oxidized to tele-methylimidazoleacetic acid. However, histamine's oxidative metabolite in the periphery, imidazoleacetic acid (IAA), is also present in brain and CSF, and its levels in brain increase after inhibition of HMT. To reinvestigate if brain has the capacity to oxidize histamine and form IAA, conscious rats were injected with [3H]histamine (10 ng), either into the lateral ventricles or cisterna magna, and decapitated 30 min later. In brains of saline-treated rats, most radioactivity recovered was due to tele-methylhistamine and tele-methylimidazoleacetic acid. However, significant amts. of tritiated IAA and its metabolites, IAA-ribotide and IAA-riboside, were consistently recovered. In rats pretreated with metoprine, an inhibitor of HMT, labeled IAA and its metabolites usually comprised the majority of histamine's tritiated metabolites. [3H]Histamine given intracisternally produced only trace amts. of oxidative metabolites. Formation of IAA, a potent GABA-A agonist with numerous neurochem. and behavioral effects, from minute quantities of histamine in brain indicates a need for reevaluation of histamine's metabolic pathway or pathways in brain and suggests a novel mechanism for interactions between histamine and the GABAergic system.

IT 2888-19-9, Imidazole-4-acetic acid-ribotide 29605-99-0, Imidazoleacetic acid-riboside

RL: BSU (Biological study, unclassified); MFM (Metabolic formation); BIOL (Biological study); FORM (Formation, nonpreparative)

(imidazoleacetic acid formation in rat brain by histamine oxidn.)

RN 2888-19-9 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-(5-0-phosphono-.beta.-D-ribofuranosyl)(9CI) (CA INDEX NAME)

RN 29605-99-0 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1983:138012 CAPLUS

DN 98:138012

TI Biliar elimination of histamine and its metabolites in guinea pigs

AU Puerta, M. L.; Ballestero, M. E. M.

CS Fac. Cienc. Biol., Univ. Complutense Madrid, Madrid, Spain

SO Comp. Biochem. Physiol. C (1983), 74C(1), 111-13 CODEN: CBPCBB; ISSN: 0306-4492

DT Journal

LA English

GΙ

$$\begin{array}{c|c} & \text{CH}_2\text{CH}_2\text{NH}_2 \\ & \text{N} & \text{I} \end{array}$$

Administration of 14C-labeled histamine (I) [51-45-6] i.v. to guinea pigs resulted in 3.5% of the radioactivity being eliminated in the bile of both males and females. Free I, methylhistamine [501-75-7], methimidazoleacetic acid [2625-49-2], imidazoleacetic acid [645-65-8] and its riboside [29605-99-0], and acetylhistamine [673-49-4] were identified in the bile. Male bile contained more free I and methylhistamine than did female bile. Evidently, biliary elimination of I and metabolites is similar to that of urine but quant. less important.

IT 29605-99-0

RL: BIOL (Biological study) (as histamine metabolite, in bile)

RN 29605-99-0 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

L5 ANSWER 6 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1974:115938 CAPLUS

DN 80:115938

TI Catabolism of orally administered carbon-14-labeled histamine in man

AU Sjaastad, Ottar; Sjaastad, O. V.

CS Inst. Surg. Res., Univ. Hosp., Oslo, Norway

SO Acta Pharmacol. Toxicol. (1974), 34(1), 33-45 CODEN: APTOA6

DT Journal

LA English

AB Within 48 hr following oral administration of 14C-labeled histamine-2HCl (I-2HCl) [56-92-8] (.sim.200 mg) to humans, 68-80% of the radioactivity was recovered in the urine, 1.8-18% was exhaled as 14CO2, and 13-19% was excreted in the feces. The main urinary I metabolites were imidazoleacetic acid [645-65-8] and methylimidazoleacetic acid [2625-49-2].

IT 29605-99-0

RL: FORM (Formation, nonpreparative)
(formation of, as histamine metabolite)

RN 29605-99-0 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2002 ACS .

AN 1971:137653 CAPLUS

DN 74:137653

TI Metabolism of [14C]-histamine in domestric animals. II. Cow and sheep

AU Eliassen, K. A.

CS Dep. Physiol., Vet. Coll. Norway, Oslo, Norway

SO Acta Physiol. Scand. (1971), 81(3), 289-99 CODEN: APSCAX

DT Journal

LA English

GI For diagram(s), see printed CA Issue.

AB In cow and sheep the oxidative deamination of histamine (I) into imidazoleacetic acid and its riboside was the major metabolic pathway. About 2% of the urinary radioactivity following 14C-labeled I injection was due to histaminol.

IT 29605-99-0

RL: FORM (Formation, nonpreparative)
 (formation of, from histamine, by ruminants)

RN 29605-99-0 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HO OH
$$CO_2H$$

L5 ANSWER 8 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1971:74587 CAPLUS

DN 74:74587

TI Uptake of [14C]-histamine by tissues of the guinea pig

AU Lewis, A. J.; Nicholls, Paul J.

CS Welsh Sch. Pharm., UWIST, Cardiff, Wales

SO J. Pharm., Pharmacol. (1971), 23(1), 66 CODEN: JPPMAB

DT Journal

LA English

GI For diagram(s), see printed CA Issue.

AB The low uptake of ring-2-14C-labeled histamine (I) (80 mg/kg, i.v.) by various tissues of guinea pigs showed that the animal, unlike cats and rabbits, does not posses an effective uptake system. The acidic metabolites of I, when detd. 8 hr after the administration, were identified as imidazole-4-acetic acid, 1-ribosylimidazole-4-acetic acid, and 1-methylimidazole-4-acetic acid.

IT 29605-99-0

RL: FORM (Formation, nonpreparative)
(formation of, from histamine by animal tissue)

RN 29605-99-0 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1969:1181 CAPLUS

DN 70:1181

TI Effects of phenethyl alcohol on yeast cells

AU Burns, Victor W.; Wong, D. L.

CS Univ. of California, David, Calif., USA

SO J. Cell. Physiol. (1968), 72(2)(Pt. 1), 97-107 CODEN: JCLLAX

DT Journal

LA English

AB The physiol. effects of phenethyl alc. (I) were studied on an exponentially growing mutant of Saccharomyces cerevisiae which required

adenine, uracil, and histidine. The incorporation of 14C-labeled glucose into the Cl3CCO2H (TCA)-sol. fraction of the cells was markedly inhibited by 0.4%, but not by 0.3% I. The incorporation of glucose into the TCA-insol. fraction was inhibited by 0.2% I. I at 0.2% reduced the turnover rate of the histidine pool and the rate of incorporation of 14Clabeled histidine into protein. The uptake of 14C-labeled adenine into the TCA-insol. fraction, an indication of RNA synthesis, was inhibited by 0.2% I. For the effect of I on DNA synthesis, 14Clabeled uracil was used as a tracer, and 0.15-0.25% I caused inhibition of the uptake of uracil. Because of a genetic block in this organism, aminoimidazole ribotide (II) accumulates in the cell in the absence of adenine. I at concns. from 0.2 to 0.6% partially prevented the accumulation of II. Gross turbidity measurements showed a decrease in cell count in the presence of I. RNA synthesis was reduced to 60% of normal at 0.15% I while DNA and protein synthesis were reduced to 90% of normal. The effects of I at <0.3% were reversible. I appears to inhibit both the substrate uptake at the cell membrane and internal RNA and DNA synthesis.

IT 27178-37-6

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (metabolism of, phenethyl alc. effect on Escherichia coli)

RN 27178-37-6 CAPLUS

CN Imidazole, amino-1-.beta.-D-ribofuranosyl- (8CI) (CA INDEX NAME)

D1-NH₂

L5 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1965:441363 CAPLUS

DN 63:41363

OREF 63:7460b-c

TI Enzymic interaction of histamine with pyridine coenzymes

AU Alivisatos, S. G. A.

CS Chicago Med. School

SO Federation Proc. (1965), Pt. 1(3), 769-73

DT Journal

LA English

The formation of histamine ribonucleoside after administration of labeled histamine was detd. in vivo. Since this compd. is most probably the in vivo degradation product of NAD, this demonstration indicates the in vivo formation of NAD. This raises the question of the potential importance of histamine-pyridine coenzyme interaction in the course of cellular metabolism in tissues rich in NADases, e.g. in the nervous tissue or during pathol. manifestation of hypersensitivity.

IT 5624-01-1, Imidazole, 4-(2-aminoethyl)-1-.beta.-D-ribofuranosyl-(formation by enzymes)

RN 5624-01-1 CAPLUS

CN 1H-Imidazole-4-ethanamine, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

L5 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1965:441136 CAPLUS

DN 63:41136

OREF 63:7422g-h

TI Evidence for the presence of imidazoleacetic acid riboside and ribotide in rat tissues

AU Robinson, Joseph D.; Green, Jack P.

CS School of Med., Yale Univ.

SO Federation Proc. (1965), 24(3;1), 777

DT Journal

LA English

AB A combination of ion-exchange and paper chromatography of the acid-sol. radioactive material from kidneys of rats given histamine-14 C showed the presence of imidazolcacetic acid riboside (I) and ribotide (II) and a third unidentified substance whose Rf value differed from all known metabolites of histamine. The most likely route for the synthesis of I and II would be oxidn. of histamine to imidazoleacetic acid followed by condensation of the acid with phosphoribosyl pyrophosphate, a reaction demonstrated in vitro; the I would then arise by dephosphorylation.

Labeled histamine adenine dinucleotide and histamine adenine dinucleotide phosphate could not be detected in kidney, liver, or brain.

IT 2888-19-9, Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl-, 5'-phosphate

(in kidneys after histamine administration)

RN 2888-19-9 CAPLUS

CN 1H-Imidazole-4-acetic acid, 1-(5-O-phosphono-.beta.-D-ribofuranosyl)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L5 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2002 ACS

AN 1964:486555 CAPLUS

DN 61:86555

OREF 61:15113g-h

TI Presence of imidazoleacetic acid riboside and ribotide in rat tissues

AU Robinson, J. D.; Green, J. P.

CS Yale Univ., School of Med., New Haven, CT

SO Nature (1964), 203(4950), 1178-9

DT Journal

LA Unavailable

AB In rats given multiple injections of **labeled** histamine (I), chromatography of trichloroacetic acid (TCA) exts. of kidney revealed 3 major radioactive fractions. These were imidazoleacetic acid riboside (II), imidazoleacetic acid ribotide (III) and an unidentified fraction not

coinciding with any of the urinary I metabolites. In brain, after injection of labeled histidine (IV), chromatography of TCA exts. revealed small fractions of total tissue radioactivity in II, III, I and imidazoleacetic acid, higher levels in unidentified metabolites, and at least 80% as IV. Radioactivity from injected I or IV was not incorporated into histamine adenine dinucleotide (HAD) or HAD phosphate in rat or

ΙT 29605-99-0, Imidazole-4-acetic acid, 1-ribosyl-RN

(in brain and kidneys after administration of histamine and histidine) 29605-99-0 CAPLUS

1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX CN

Absolute stereochemistry.

L5 ANSWER 13 OF 15 CAPLUS COPYRIGHT 2002 ACS

1964:456802 CAPLUS ΑN

DN 61:56802

OREF 61:9880f-h,9881a

Increased turnover of phosphoribosylpyrophosphate, a purine nucleotide TIprecursor, in certain gouty subjects

ΑU Wyngaarden, J. B.; Jones, O. W.; Ashton, D. M.

Atti Congr. Lega Intern. Reumatismo, 10.degree., Rome (1961), 1, 249-53 SO DT

LA

Since phosphoribosylpyrophosphate (I) is an obligatory precursor of purine AB nucleotides, its turnover has been investigated in gouty subjects. The hyperuricemia of gout may be due to overproduction or underexcretion of uric acid, or both. Orally administered imidazoleacetic acid (II) is partially excreted in urine as the imidazoleacetic acid ribonucleotide (III), and if glucose-14C is given simultaneously the ribose moiety is labeled. It is assumed that the same "pool" of I is involved both in the production of III and of phosphoribosylamine (the 1st specific precursor of purine nucleotides). Subjects were all males. Five controls had no gout or renal disease personally, or in the family history. Five gouty patients varied from asymptomatic hyperuricemia to advanced chronic tophaceous gout. All were given 25 .mu.c. glucose-U-14C and 20 micromoles/kg. II. Urine was collected in 5 ml. of concd. HCl, either in 2-hr. aliquots, or in a single 10-hr. sample, and stored at 4.degree... CO2 was then removed by aeration, the pH adjusted to 8, the III collected on a Dowex-1 (acetate) column, and purified on a Dowex-50 (H+) column. The product in M citrate, pH 6.0, was hydrolyzed with a bacterial riboside hydrolase, and the protein-free filtrate passed through a mixed-bed resin (MB-3, Fisher), and the eluate analyzed for 14C and ribose (orcinol). Uric acid was detd. by differential spectrophotometry using uricase. In the controls, 0.010-0.047% 14C was incorporated into urinary III in 10 hours. For 2 gouty subjects with low and normal uric acid excretions, the corresponding figures were 0.009 and 0.058%, and for 3 gouty hyperuricemic subjects the range was 0.1640.309%. In these latter 3 subjects, the sp. activity (counts/ min./mg.) of the ribose moiety of III was approx. 8 times that of the controls. If urine were collected in 2-hr. aliquots, the max. sp. activity occurred about 2 hrs. earlier in all gouty subjects than in controls, and the peak values for the 3 hyperexcretors were 2-4-fold greater than controls. There was an increased I turnover in the

3 hyperexcretor gouty subjects, but there may be a continuous gradation in

2888-19-9, Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl-, ΙT 5'-phosphate (in urine in gout)

RN 2888-19-9 CAPLUS

1H-Imidazole-4-acetic acid, 1-(5-0-phosphono-.beta.-D-ribofuranosyl)-CN

Absolute stereochemistry.

ANSWER 14 OF 15 CAPLUS COPYRIGHT 2002 ACS L5 AN

1963:470932 CAPLUS

DN 59:70932

OREF 59:13187f-g

In vivo degradation of histamine-adenine dinucleotide phosphate to ΤI histamine ribonucleoside ΑU

Alivisatos, S. G. A.; Abdel-Latif, A. A.; Ungar, F.; Mourkides, G. A. CS

SO Nature (1963), 199(4896), 907-8 DΤ

Journal

LΑ Unavailable

Mice were injected intraperitoneally with a histamine-adenine dinucleotide AB phosphate. The histamine was labeled with C14 at C-2. The only radioactive excretion in the urine of mice was identified as histamine IT

5624-01-1, Imidazole, 4-(2-aminoethyl)-1-.beta.-D-ribofuranosyl-(formation from histamine-adenine dinucleotide phosphate in vivo) RN5624-01-1 CAPLUS

1H-Imidazole-4-ethanamine, 1-.beta.~D-ribofuranosyl- (9CI) CN NAME) (CA INDEX

Absolute stereochemistry.

ANSWER 15 OF 15 CAPLUS COPYRIGHT 2002 ACS L5 AN

1962:74743 CAPLUS

DN 56:74743

OREF 56:14584b-d

The catabolism of tissue nucleic acid. III. The catabolism of ribonucleic ΤI acid after total-body x-irradiation Gerber, Georg B.; Gerber, Gisela; Altman, Kurt I.

ΑIJ CS

Univ. of Rochester, Rochester, NY

SO Intern. J. Radiation Biol. (1961), 4, 67-73

DT

LΑ Unavailable

cf. CA 54, 25159h. The effect of total-body x-irradiation on ribonucleic AΒ acid (RNA) catabolism was studied in rats whose RNA had been labeled by injection of glucose-U-C14 three days previously. sp. activity of urinary ribosyl imidazole acetate (I) as well as of RNA of liver, intestine, muscle, spleen and thymus was detd. after x- or sham-irradiation. Rats were either pair fed or starved after irradiation. After 1000 r. the sp. activity of I was increased whereas that of intestinal and muscle RNA was decreased with little change in liver RNA. In pair-fed animals, exposure to 756 r. decreased the sp. activity of spleen and thymus RNA, and gave a steady decrease in that of I over 5 days. Sp. activity of liver RNA was diminished in the sham-irradiated rats and was equiv. to that of I which was excreted on the first day after treatment. It was concluded that radiation-induced increase in RNA catabolism is present mainly in intestine and muscle on the second and third day after exposure whereas starvation-induced increase in catabolism occurs primarily in the liver and on the first day of starvation. IT 29605-99-0, Imidazole-4-acetic acid, 1-ribosyl-

(in urine after x-ray irradiation)

RN29605-99-0 CAPLUS

1H-Imidazole-4-acetic acid, 1-.beta.-D-ribofuranosyl- (9CI) CN (CA INDEX

Absolute stereochemistry.

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FILE 'HOME' ENTERED AT 06:49:39 ON 25 SEP 2002

=> file reg

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

0.21

0.21

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 23 SEP 2002 HIGHEST RN 454421-17-1 DICTIONARY FILE UPDATES: 23 SEP 2002 HIGHEST RN 454421-17-1

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

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Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

*** YOU HAVE NEW MAIL ***

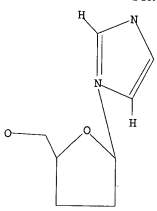
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Uploading 09880727.str

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

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SEARCH TIME: 00.00.03

7 ANSWERS

L2 7 SEA SSS FUL L1

=> file caplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 140.28 140.49

FULL ESTIMATED COST

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FILE COVERS 1907 - 25 Sep 2002 VOL 137 ISS 13 FILE LAST UPDATED: 23 Sep 2002 (20020923/ED)

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=> s 12

L3 3 L2

=> d l3 bib abs hitstr 1-3

L3 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS

AN 2000:669931 CAPLUS

DN 134:5110

TI Contribution of the Adenine Base to the Activity of Adenophostin A Investigated Using a Base Replacement Strategy

AU Marwood, Rachel D.; Jenkins, David J.; Correa, Vanessa; Taylor, Colin W.; Potter, Barry V. L.

CS Wolfson Laboratory of Medicinal Chemistry Department of Pharmacy and Pharmacology, University of Bath, Bath, BA2 7AY, UK

Journal of Medicinal Chemistry (2000), 43(22), 4278-4287 CODEN: JMCMAR; ISSN: 0022-2623

PB American Chemical Society

DT Journal

LA English

AB Syntheses of 3'-O-.alpha.-D-glucopyranosyl-1-.beta.-D-ribofuranosidoimidazole 2',3'',4''-trisphosphate (I) and 3'-O-.alpha.-D-glucopyranosyl-9-.beta.-D-ribofuranosidopurine 2',3'',4''-trisphosphate (II), two analogs of the superpotent 1D-myo-inositol 1,4,5-trisphosphate receptor agonist adenophostin A (III), are described. 5-O-Benzyl-1,2-O-isopropylidene-.alpha.-D-ribofuranose was prepd. by an improved route from 1,2-O-isopropylidene-.alpha.-D-

xylofuranose and was coupled with 3,4-di-O-acetyl-2,6-di-O-benzyl-Dglucopyranosyl di-Me phosphite to give 3',4'-di-O-acetyl-2',5,6'-tri-Obenzyl-3-0-.alpha.-D-glucopyranosyl-1,2-0-isopropylidene-.alpha.-Dribofuranose. Removal of the isopropylidene acetal and subsequent acetylation gave the central disaccharide 1,2,3',4'-tetra-O-acetyl-2',5,6'tri-O-benzyl-3-O-.alpha.-D-glucopyranosyl-D-ribofuranose. Vorbuggen condensation with activated imidazole or purine gave the required .beta.-substituted derivs. which were further elaborated to I and II, resp. Radioligand binding assays to hepatic InsP3 receptors and functional assays of Ca2+ release from permeabilized hepatocytes gave a rank order of potency of the ligands III .apprxeq. II > I .apprxeq. Ins(1,4,5)P3 indicating that the N6-amino group of III is of little importance for activity and that a min. of a two-fused-ring nucleobase is required for activity to exceed that of Ins(1,4,5)P3. The role of the adenine base in the activity of the adenophostins is discussed. general method should facilitate ready access to nucleobase-modified adenophostin analogs for SAR studies.

IT 308290-83-7P

CN

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and membrane binding structure activity relationships of Adenophostin A via a coupling reaction)

RN 308290-83-7 CAPLUS

1H-Imidazolium, 1,3-bis[2-O-acetyl-3-O-[3,4-di-O-acetyl-2,6-bis-O-(phenylmethyl)-.alpha.-D-glucopyranosyl]-5-O-(phenylmethyl)-.beta.-D-ribofuranosyl]- (9CI) (CA INDEX NAME)

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RE.CNT 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS

AN 1996:762145 CAPLUS

DN 126:144489

TI New C2 symmetrical and semi-symmetrical substituted imidazolium ribonucleoside. Imidazolic nucleosides analogs

AU Mourabit, A. Al; Beckmann, M.; Poupat, C.; Ahond, A.; Potier, P.

CS Institut de Chimie des Substances Naturelles du C.N.R.S., Gif-sur-Yvette, 91198, Fr.

SO Tetrahedron: Asymmetry (1996), 7(12), 3455-3464 CODEN: TASYE3; ISSN: 0957-4166

Ι

PB Elsevier

DT Journal

LA English

GI

AB New C2 sym. imidazolium nucleosides, e.g. I, have been synthesized using silyl Hilbert Johnson-Vorbrugen method and coupling of trimethylsilylimidazole with the peracylated D-ribofuranose. The was used. These new base modified nucleosides were devoid of activity against HIV and cytotoxicity.

IT 186648-75-9P 186648-80-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(prepn. of C2 sym. and semi-sym. substituted imidazolium ribonucleosides as virucides)

RN 186648-75-9 CAPLUS

CN 1H-Imidazolium, 1,3-bis(2,3,5-tri-O-acetyl-.beta.-D-ribofuranosyl)-, hydroxide (9CI) (CA INDEX NAME)

OH-

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 186648-80-6 CAPLUS

CN 1H-Imidazolium, 4-[(1S,2S)-3-azido-2-chloro-1-[[dimethyl(1,1,2-trimethylpropyl)silyl]oxy]propyl]-1,3-bis(2,3,5-tri-O-benzoyl-.beta.-D-ribofuranosyl)-, hydroxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

IT 186648-73-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of C2 sym. and semi-sym. substituted imidazolium ribonucleosides as virucides)

RN 186648-73-7 CAPLUS

CN 1H-Imidazolium, 1,3-bis(2,3,5-tri-O-benzoyl-.beta.-D-ribofuranosyl)-, hydroxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

IT 186648-77-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of C2 sym. and semi-sym. substituted imidazolium ribonucleosides as virucides)

RN 186648-77-1 CAPLUS

CN 1H-Imidazolium, 1,3-di-.beta.-D-ribofuranosyl-, hydroxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

OH-

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

L3 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS

AN 1992:470204 CAPLUS

DN 117:70204

TI Nucleosides. 163. Synthesis of ribosides and ribotides of imidazole-4(5)-acetic acid and 1-methylimidazole-4(5)-acetic acid

AU Matulic-Adamic, Jasenka; Watanabe, Kyoichi A.

CS Lab. Org. Chem., Sloan-Kettering Inst. Cancer Res., New York, NY, 10021, USA

SO Korean J. Med. Chem. (1991), 1(1), 54-64. CODEN: KJMCE7

DT Journal

LA English

OS CASREACT 117:70204

GI

AB Nucleotide imidazoleacetic acid, e.g. I, were prepd. from imidazole-4(5)-acetonitrile (II). Regioselective tritylation of II followed by N-methylation with Me2S and hydrolysis gave 1-methylimidazole-5-acetic acid.

IT 142527-55-7P 142606-76-6P

Ι

RN 142527-55-7 CAPLUS

CN 1H-Imidazolium, 4-(carboxymethyl)-3-methyl-1-.beta.-D-ribofuranosyl-, inner salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 142606-76-6 CAPLUS

CN 1H-Imidazolium, 4-(carboxymethyl)-3-methyl-1-(5-O-phosphono-.beta.-D-ribofuranosyl)-, inner salt, monosodium salt (9CI) (CA INDEX NAME)

$$H_2O_3PO-CH_2$$
 O N N Me HO OH $CH_2-CO_2-CH_2$

Na

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

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=> file reg COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

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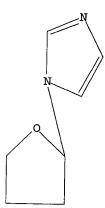
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*** YOU HAVE NEW MAIL ***

=> Uploading 09880727.str

L1 STRUCTURE UPLOADED

=> d l1 L1 HAS NO ANSWERS L1 STR



Structure attributes must be viewed using STN Express query preparation.

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100.0% PROCESSED 108494 ITERATIONS SEARCH TIME: 00.00.04

104846 ANSWERS

L2 104846 SEA SSS FUL L1

=> file caplus
COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 140.28 140.49

FULL ESTIMATED COST

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=> s 12

L3 259099 L2

=> s l3 and label

48825 LABEL 4 2754 L3 AND LABEL

=> s 14 and coupled label?

234873 COUPLED 374492 LABEL?

4 COUPLED LABEL?

(COUPLED (W) LABEL?)

L5 0 L4 AND COUPLED LABEL?

=> d l4 bib abs hitstr 1, 200, 2754

L4 ANSWER 1 OF 2754 CAPLUS COPYRIGHT 2002 ACS

AN 2002:688459 CAPLUS

DN 137:180760

TI DNA sequencing using multiple fluorescent labels being distinguishable by their decay times

IN Jensen, Morten; Parce, J. Wallace

PA Caliper Technologies Corp., USA

SO U.S., 18 pp., Cont.-in-part of U.S. Ser. No. 132,191, abandoned. CODEN: USXXAM

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DΤ
     Patent
LA
     English
FAN.CNT 2
     PATENT NO.
                      KIND DATE
                                         APPLICATION NO.
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ΡI
     US 6447724
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                                                          19990811
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     EP 1104491
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             IE, FI
PRAI US 1998-132191
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     US 1998-122064P
                      P
                           19980811
     US 1998-132181
                      Α
                           19980811
    US 1998-132554
                      A
                           19980811
    US 1998-213297
                      Α
                           19981215
    WO 1999-US18294
                    W
                           19990811
AB
```

AB A method is provided for identifying components of a mixt. by labeling the individual components with fluorescent agents having different fluorescence lifetimes. The components are subsequently sepd., fluorescent labels detected and their lifetimes measured. Based on the measured fluorescent lifetimes, the components of mixts. of small org. identified.

IT 39007-51-7, Ethenoadenosine
RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
(Analytical study); BIOL (Biological study); USES (Uses)
(DNA sequencing using multiple fluorescent labels being distinguishable
by their decay times)

RN 39007-51-7 CAPLUS

CN 3H-Imidazo[2,1-i]purine, 3-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
AN 2000:402045 CAPLUS
DN 133:40226
TI Targeted molecular bar codes and methods for using the same
IN Akeson, Mark; Deamer, David W.; Vercoutere, Wenonah; Olsen, Hugh E.;
Braslau, Rebecca; Singaram, Bakthan; Steiner, Derek; Cappuccio, Frank
PA The Regents of the University of California, USA
PCT Int. Appl., 53 pp.
```

ANSWER 200 OF 2754 CAPLUS COPYRIGHT 2002 ACS

CODEN: PIXXD2

DT Patent LA English FAN.CNT 1

L4 AN

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 1999-US30598 19991210 WO 2000034527 20000615 **A2** ΡI 20001012 WO 2000034527 A3 AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, W: CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG PRAI US 1998-111802P 19981211 P P 19991006 US 1999-158020P Targeted mol. bar codes and methods for using the same are provided. The AΒ

Targeted mol. bar codes and methods for using the same are provided. The subject targeted mol. bar codes include a mol. bar code and a member of a specific binding pair, where the specific binding pair member is generally bonded to the bar code through a linking group. The subject mol. bar code may be read during translocation through a single nano-meter scale pore. The subject targeted mol. bar codes find use in a variety of different applications involving analyte detection, such as screening and diagnostic applications. A targeted bar code composed of poly dT18 linked by a disulfide bond to a 50 base-long antisense segment of N-ras Exon 1 was synthesized and used to detect N-ras in a mixt. The target oligonucleotides were attached to polystyrene beads. Following hybridization and recovery and washing of the bound bar code, the disulfide linkage was cleaved, the beads were spun down, and the bar code-contg. supernatant was added to an .alpha.-hemolysin nanopore miniature device for detection.

(in prepn. of mol. bar codes; targeted mol. bar codes and methods for using same)

RN 24937-83-5 CAPLUS

CN 5'-Adenylic acid, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 61-19-8

CMF C10 H14 N5 O7 P

Absolute stereochemistry.

L4 ANSWER 2754 OF 2754 CAPLUS COPYRIGHT 2002 ACS

AN 1962:14648 CAPLUS

DN 56:14648

OREF 56:2772i,2773a-b

TI The effect of puromycin on protein metabolism and cell division in fertilized sea urchin eggs

AU Hultin, T.

CS WennerGren Inst., Stockholm

SO Experientia (1961), 17, 410-11

DT Journal LA English

Puromycin introduced into sea urchin eggs just before fertilization at concns. above 10-4M halts division at the "clear streak" stage. A correlation was sought between this effect and the inhibitory action of puromycin on the incorporation of labeled amino acids into protein in the eggs. L-Valine-C14 was used as the label. The inhibition of division in labeled whole eggs varied with the concn. of puromycin. At a concn. of 10-4M, 50% stopped at the "clear streak" stage, while at 10-5M 40% reached a 4-cell stage. The cell-free incorporation system showed a greater sensitivity to puromycin. The mitotic block is probably a direct effect of impaired protein metabolism.

IT 5682-30-4, Adenosine, 3'-(.alpha.-amino-p-methoxyhydrocinnamamido)-3'-deoxy-N,N-dimethyl-

(effect on cell division and protein metabolism in sea urchin ova)

RN 5682-30-4 CAPLUS

CN Adenosine, 3'-[[2-amino-3-(4-methoxyphenyl)-1-oxopropyl]amino]-3'-deoxy-N,N-dimethyl-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

=> d his

(FILE 'HOME' ENTERED AT 15:40:21 ON 24 SEP 2002)

FILE 'REGISTRY' ENTERED AT 15:40:38 ON 24 SEP 2002

L1 STRUCTURE UPLOADED

L2 104846 S L1 FULL

FILE 'CAPLUS' ENTERED AT 15:41:18 ON 24 SEP 2002

L3 259099 S L2

L4 2754 S L3 AND LABEL

L5 0 S L4 AND COUPLED LABEL?

=> s 14 and imidazole

42254 IMIDAZOLE

L6 23 L4 AND IMIDAZOLE

=> d 16 bib abs hitstr 1-23

L6 ANSWER 1 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 2000:748405 CAPLUS

DN 134:42020

TI Carbon-14 labeling of a potential new immunoregulant agent

AU Egan, M. A. McEvoy; Dean, D. C.; Marks, T. M.; Song, Zhiguo; Melillo, D.

G.

CS Merck Research Laboratories, Rahway, NJ, 07065, USA

SO Journal of Labelled Compounds & Radiopharmaceuticals (2000), 43(11), 1095-1105

CODEN: JLCRD4; ISSN: 0362-4803

PB John Wiley & Sons Ltd.

DT Journal

LA English

OS CASREACT 134:42020

GΙ

RN

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB A carbon-14 labeled version of the ascomycin analog I, a potential new immunosuppressant agent, was synthesized for utilization in animal and human drug metab. studies. In order to place the carbon-14 label at a metabolically stable position, it was necessary to modify the established synthesis of a key intermediate. [14C]-I is prepd. by a highly chemoselective alkylation of ascomycin at the C-32 hydroxy position with a carbon-14 labeled imidazolyl trichloroimidate side chain II. Carbon-14 was efficiently incorporated in II through carboxylation of an imidazole C-2 lithiate with [14C] carbon dioxide.

IT 312583-35-0

RL: RCT (Reactant); RACT (Reactant or reagent)
(carbon-14 labeling of ascomycin as a potential new immunoregulant

agent ascomycin) 312583-35-0 CAPLUS

CN 1H-Imidazole-2-carboxylic-14C acid, 4-(3,5-dimethoxyphenyl)-1-(tetrahydro-2-furanyl)-, lithium salt (9CI) (CA INDEX NAME)

• Li

IT 312583-34-9P 312583-36-1P 312583-38-3P 312583-39-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(carbon-14 labeling of ascomycin as a potential new immunoregulant agent ascomycin)

RN 312583-34-9 CAPLUS

CN Ethanimidic acid, 2,2,2-trichloro-, [4-(3,5-dimethoxyphenyl)-1-(tetrahydro-2-furanyl)-1H-imidazol-2-yl]methyl-14C ester (9CI) (CA INDEX NAME)

RN 312583-36-1 CAPLUS

CN 1H-Imidazole, 4-(3,5-dimethoxyphenyl)-1-(tetrahydro-2-furanyl)- (9CI) (CA INDEX NAME)

RN 312583-38-3 CAPLUS

CN 1H-Imidazole-2-carboxylic-14C acid, 4-(3,5-dimethoxyphenyl)-1-(tetrahydro-2-furanyl)-, methyl ester (9CI) (CA INDEX NAME)

RN 312583-39-4 CAPLUS

CN 1H-Imidazole-2-methanol-.alpha.-14C, 4-(3,5-dimethoxyphenyl)-1-(tetrahydro-2-furanyl)- (9CI) (CA INDEX NAME)

RE.CNT 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1999:491591 CAPLUS

DN 131:257779

TI 15N-Multi-labeled Adenine and Guanine Nucleosides. Syntheses of [1,3,NH2-15N3] - and [2-13C-1,3,NH2-15N3] - Labeled Adenosine, Guanosine, 2'-Deoxyadenosine, and 2'-Deoxyguanosine

AU Abad, Jose-Luis; Gaffney, Barbara L.; Jones, Roger A.

CS Department of Chemistry, Rutgers The State University of New Jersey, Piscataway, NJ, 08854, USA

SO Journal of Organic Chemistry (1999), 64(18), 6575-6582 CODEN: JOCEAH; ISSN: 0022-3263

PB American Chemical Society

DT Journal

LA English

AB The authors report a high-yield route to the following specifically 15Nand 13C-multi-labeled nucleosides: [1,3,NH2-15N3] - and [2-13C-1,3,NH2-15N3]-adenosine; [1,3,NH2-15N3]- and [2-13C-1,3,NH2-15N3]guanosine; [1,3,NH2-15N3] - and [2-13C-1,3,NH2-15N3]-2'-deoxyadenosine; [1,3,NH2-15N3] - and [2-13C-1,3,NH2-15N3]-2'-deoxyguanosine. In each set, the 13C2 atom functions as a "tag" that allows the 15N1 and 15N3 atoms to be unambiguously differentiated from the untagged versions in 15N NMR of RNA or DNA fragments. The key intermediate of this synthetic strategy for both the adenine and guanine nucleosides is [NH2,CONH2-15N2]-5-amino-4imidazolecarboxamide. The [2-13C]-label is added through a ring closure using [13C]-sodium Et xanthate (NaS13CSOEt). Enzymic transglycosylation of either multi-labeled 6-chloropurine or multi-labeled 2-mercaptohypoxanthine and a final reaction with 15NH3 give the adenine and guanine nucleosides. This is the first report of a [3-15N]-labeled guanine nucleoside.

IT 244769-65-1P 244769-69-5P 244769-73-1P 244769-74-2P 244769-75-3P 244769-81-1P 244769-82-2P 244769-83-3P

RL: BPN (Biosynthetic preparation); RCT (Reactant); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent) (syntheses of [15N3] - and [13C,15N3] - labeled adenosine, guanosine, deoxyadenosine, and deoxyguanosine nucleosides)

RN 244769-65-1 CAPLUS

CN 9H-Purine-1,3-15N2, 6-chloro-9-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-69-5 CAPLUS

CN Xanthosine-1,3-15N2, 2-S-methyl-2-thio- (9CI) (CA INDEX NAME)

RN 244769-73-1 CAPLUS

CN 9H-Purine-2-13C-1,3-15N2, 6-chloro-9-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-74-2 CAPLUS

CN 9H-Purine-1,3-15N2, 6-chloro-9-(2-deoxy-.beta.-D-erythro-pentofuranosyl)(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-75-3 CAPLUS

CN 9H-Purine-2-13C-1,3-15N2, 6-chloro-9-(2-deoxy-.beta.-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

244769-81-1 CAPLUS RNCNXanthosine-2-13C-1,3-15N2, 2-S-methyl-2-thio- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN244769-82-2 CAPLUS

Xanthosine-1,3-15N2, 2'-deoxy-2-S-methyl-2-thio- (9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

$$\begin{array}{c|c} & & & \\ & & & \\ 15_N & & & \\ & & & \\ Mes & & \\ 15_N & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

RN244769-83-3 CAPLUS

Xanthosine-2-13C-1,3-15N2, 2'-deoxy-2-S-methyl-2-thio- (9CI) (CA INDEX CN NAME)

IT 118-00-3, Guanosine, reactions 961-07-9,

2'-Deoxyguanosine

RL: RCT (Reactant); RACT (Reactant or reagent)

(syntheses of [15N3] - and [13C,15N3] - labeled adenosine, guanosine, deoxyadenosine, and deoxyguanosine nucleosides)

RN 118-00-3 CAPLUS

CN Guanosine (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 961-07-9 CAPLUS

CN Guanosine, 2'-deoxy- (6CI, 8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 20244-86-4P, 7-Methylguanosine 244769-70-8P

244769-84-4P 244769-85-5P 244769-86-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(syntheses of [15N3] - and [13C,15N3] - labeled adenosine, guanosine, deoxyadenosine, and deoxyguanosine nucleosides)

RN 20244-86-4 CAPLUS

CN 1H-Purinium, 2-amino-6,9-dihydro-7-methyl-6-oxo-9-.beta.-D-ribofuranosyl-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 244769-70-8 CAPLUS

CN Inosine-1,3-15N2, 2-(methylsulfinyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-84-4 CAPLUS

CN Inosine-2-13C-1,3-15N2, 2-(methylsulfinyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-85-5 CAPLUS

CN Inosine-1,3-15N2, 2'-deoxy-2-(methylsulfinyl)- (9CI) (CA INDEX NAME)

Me
$$15_N$$
 N R R R O OH

RN 244769-86-6 CAPLUS

CN Inosine-2-13C-1,3-15N2, 2'-deoxy-2-(methylsulfinyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

$$15_{\mathrm{NH}_2}$$
 15_{NH_2} $15_{$

RN 244769-66-2 CAPLUS CN Adenosine-N,1,3-15N3 (9CI) (CA INDEX NAME)

RN 244769-76-4 CAPLUS CN Adenosine-2-13C-N,1,3-15N3 (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-77-5 CAPLUS CN Adenosine-N,1,3-15N3, 2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-78-6 CAPLUS CN Adenosine-2-13C-N,1,3-15N3, 2'-deoxy- (9CI) (CA INDEX NAME)

RN 244769-87-7 CAPLUS CN Guanosine-2-13C-N,1,3-15N3 (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-88-8 CAPLUS CN Guanosine-N,1,3-15N3, 2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 244769-89-9 CAPLUS CN Guanosine-2-13C-N,1,3-15N3, 2'-deoxy- (9CI) (CA INDEX NAME)

RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1999:143185 CAPLUS

DN 130:308594

TI Preparation of an IMI dye (imidazole functional group) containing a 4-(N,N-dimethylaminosulfonyl)-2,1,3-benzoxadiazole fluorophore for labeling of phosphomonoesters

AU Lan, Zhang-Hua; Qian, Xiaohua; Giese, Roger W.

CS Department of Pharmaceutical Sciences in the Bouve College of Pharmacy and Health Professions, Barnett Institute, and Chemistry Department, Northeastern University, Boston, MA, 02115, USA

SO Journal of Chromatography, A (1999), 831(2), 325-330 CODEN: JCRAEY; ISSN: 0021-9673

PB Elsevier Science B.V.

DT Journal

LA English

We are studying dye-imidazole conjugates ("IMI dyes") as reagents for labeling phosphomonoesters such as nucleotides. Previously we have employed a BODIPY dye in our IMI reagents, and analyzed the labeled products by capillary electrophoresis with laser-induced fluorescence detection (CE-LIF) involving an argon ion laser. (The BODIPY fluorophore is a 4,4-difluoro-4-bora-3a,4a-diaza-s-indacene). Here we broaden the technol. by prepg. a DBD-IMI dye [DBD = 4-(N,N-dimethylaminosulfonyl)-2,1,3-benzoxadiazole], and using a helium-cadmium laser. While DBD-IMI (IMI3) is about 50x more stable photolytically than a BODIPY-IMI dye (IMI2, a conjugate of a BODIPY dye with histamine, was tested), the detection limit for IMI2 (5.cntdot.10-11 M; SIN=5, CE-LIF with an argon ion laser) is tenfold better than that for IMI3 (5.cntdot.10-10 M, SIN=5, helium-cadmium laser). IMI3 conjugates of the four major DNA nucleotides were prepd. and detected by CE-LIF.

IT 61-19-8, 5'-AMP, analysis 85-32-5, 5'-GMP
RL: ANT (Analyte); BSU (Biological study, unclassified); RCT (Reactant);
ANST (Analytical study); BIOL (Biological study); RACT (Reactant or reagent)

(prepn. of an IMI dye (imidazole functional group) contg. a 4-(N,N-dimethylaminosulfonyl)-2,1,3-benzoxadiazole fluorophore for labeling of phosphomonoesters)

RN 61-19-8 CAPLUS

CN 5'-Adenylic acid (8CI, 9CI) (CA INDEX NAME)

RN 85-32-5 CAPLUS

CN 5'-Guanylic acid (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$H_2N$$
 H_2N
 H_3
 H_4
 H_5
 H_6
 H_6
 H_7
 H_8
 H_8
 H_9
 $H_$

RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1998:343335 CAPLUS

DN 129:51578

TI Phosphate-specific fluorescence labeling with BO-IMI: reaction details

AU Wang, Poguang; Giese, Roger W.

CS Chemistry Department, Barnett Institute, Department of Pharmaceutical Sciences in the Bouve College of Pharmacy and Health Professions, Northeastern University, Boston, MA, 02115, USA

SO Journal of Chromatography, A (1998), 809(1 + 2), 211-218 CODEN: JCRAEY; ISSN: 0021-9673

PB Elsevier Science B.V.

DT Journal

LA English

Previously the authors reported that BO-IMI, a reagent which contains a BODIPY fluorophore linked to an imidazole group, can be used to covalently label a phosphomonoester in a single step under aq. conditions [P. Wang, R.W. Giese, Anal. Chem . 65(1993) 3518]. reaction was conducted in the presence of a water-sol. carbodiimide 1-ethyl-3-(3'-N,N'-dimethylaminopropyl)carbodiimide [EDC] to activate the phosphomonoester, and the coupling took place onto both the N1 and N3 imidazole nitrogens of BO-IMI. Whether the two BO-IMI-phosphomonoester regioisomers migrated sep. or together during capillary electrophoresis depended on the pH, due to a difference in their pKa values. Since then, the authors have studied the reaction in more detail leading to the information reported here. First, the regioisomer ratio changes during the reaction, and found that the mechanism involves both spontaneous and BO-IMI-catalyzed hydrolysis of the less stable isomer. Second, there is a background reaction in which BO-IMI becomes attached to EDC. Third, the BO-IMI-phosphomonoester product (a mixt. of two isomers), that is obsd. by capillary electrophoresis at an alk. pH, is found to no longer contain the two fluorine atoms present in the starting BO-IMI reagent. This is because they are replaced by hydroxy groups at

high pH. Finally, an event was discovered which complicates the detection of .ltorsim.60 fmol of a phosphomonoester with BO-IMI: hydrolysis of a tiny fraction of the BO-IMI takes place during the coupling reaction, which leads to chem. noise in the capillary electropherogram.

IT 208529-99-1

RL: ANT (Analyte); FMU (Formation, unclassified); ANST (Analytical study); FORM (Formation, nonpreparative) (capillary electrophoresis of)

RN 208529-99-1 CAPLUS

CN Borate(1-), [N-acetyl-1-(2'-deoxy-7,8-dihydro-8-oxo-5'-adenylyl)-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

● H+

IT 208529-95-7 208529-97-9 208530-05-6 208530-07-8 208530-09-0

RL: ANT (Analyte); FMU (Formation, unclassified); ANST (Analytical study); FORM (Formation, nonpreparative)

(formation in phosphate-specific fluorescence labeling with BO-IMI)

RN 208529-95-7 CAPLUS

CN Borate(1-), [N-acetyl-1-[2'-deoxy-8-(phenylmethoxy)-5'-adenylyl]-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-A

● H+

PAGE 2-B

 \sim O- CH₂- Ph

RN 208529-97-9 CAPLUS

CN Borate(1-), [N-acetyl-1-[8-(acetyl-9H-fluoren-2-ylamino)-2'-deoxy-5'-guanylyl]-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-A

● H+

PAGE 2-B

RN 208530-05-6 CAPLUS

CN Borate(1-), [N-acetyl-3-[2'-deoxy-8-(phenylmethoxy)-5'-adenylyl]-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

RN 208530-07-8 CAPLUS
CN Borate(1-), [N-acetyl-3-[8-(acetyl-9H-fluoren-2-ylamino)-2'-deoxy-5'-guanylyl]-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-B

 \sim NH₂

● H+

RN 208530-09-0 CAPLUS
CN Borate(1-), [N-acetyl-3-(2'-deoxy-7,8-dihydro-8-oxo-5'-adenylyl)-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-B

● H+

Borate(1-), [N-acetyl-1-(2'-deoxy-5'-adenylyl)-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]difluoro-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-B

— ОН

RN 208529-79-7 CAPLUS

CN Borate(1-), [N-acetyl-3-(2'-deoxy-5'-adenylyl)-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]difluoro-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

● H+

RN 208529-83-3 CAPLUS

CN Borate(1-), [N-acetyl-1-(2'-deoxy-5'-adenylyl)-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-B

— ОН

RN 208529-85-5 CAPLUS

CN Borate(1-), [N-acetyl-3-(2'-deoxy-5'-adenylyl)-L-histidine 2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(2-)]dihydroxy-, hydrogen, (T-4)- (9CI) (CA INDEX NAME)

PAGE 1-A

Me
$$N_{3+N}$$
 CH_{2} CH_{2}

PAGE 1-B

● H+

RN 208530-03-4 CAPLUS

CN Borate(8-), [N-acetyl-1-(2'-deoxyguanylyl-(5'.fwdarw.3')-thymidylyl-(5'.fwdarw.3')-thymidylyl-(5'.fwdarw.3')-2'-deoxycytidylyl-(5'.fwdarw.3')-2'-deoxyguanylyl-(5'.fwdarw.3')-2'-deoxyadenylyl-(5'.fwdarw.3')-2'-deoxyadenylyl-(5'.fwdarw.3')-2'-deoxyadenylyl-L-histidine
2-[3-[5-[(3,5-dimethyl-2H-pyrrol-2-ylidene-.kappa.N)methyl]-1H-pyrrol-2-yl-.kappa.N]-1-oxopropyl]hydrazidato(9-)]dihydroxy-, octahydrogen, (T-4)-(9CI) (CA INDEX NAME)

PAGE 1-B

PAGE 3-A

●8 H+

PAGE 3-B

NH₂

IT 653-63-4

RL: ANT (Analyte); ANST (Analytical study)

(phosphate-specific fluorescence labeling with BO-IMI)

RN 653-63-4 CAPLUS

CN 5'-Adenylic acid, 2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

IT 902-04-5

RL: ANT (Analyte); ANST (Analytical study)
(phosphate-specific fluorescence labeling with BO-IMI and capillary electrophoresis of)

RN 902-04-5 CAPLUS

CN 5'-Guanylic acid, 2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$H_2N$$
 N
 N
 R
 R
 R
 R
 OPO_3H_2
 OH

L6 ANSWER 5 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1997:653036 CAPLUS

DN 127:328923

TI Ammonium assimilation in bryophytes. L-Glutamine synthetase from Sphagnum fallax

AU Kahl, Stefan; Gerendas, Joska; Heeschen, Volker; Ratcliffe, R. George; Rudolph, Hansjorg

CS Biologiezentrum, Botanisches Inst. der Christian-Albrechts-Univ. zu Kiel, Kiel, D-24098, Germany

SO Physiologia Plantarum (1997), 101(1), 86-92 CODEN: PHPLAI; ISSN: 0031-9317

PB Munksqaard

DT Journal

LA English

AΒ Cytosolic and plastidic L-glutamine synthetase (E.C. 6.3.1.2) isoenzymes from Sphagnum fallax Klinggr. (Klinggr. clone 1) were sepd. by size-exclusion and ion exchange chromatog. The cytosolic enzyme (GS1) was purified to apparent electrophoretic homogeneity. The native enzyme had a mol. mass of 390 .+-. 20 kDa as estd. by gel filtration and was apparently composed of 8 subunits with mol. masses of 48 kDa. GS1 activity could be measured from pH 6.8 to 8.6 in 50 mM imidazole buffer, with a broad optimum between pH 7.2 and 8.0. The Km values were 2.5, 0.5, and 0.5 mM for L-glutamate, ammonium, and ATP, resp. The enzyme was inhibited by more than 10 mM ammonium or glutamate. The incorporation of 15NH4+into amino acids was obsd. in vivo using 15N NMR. Label from ammonium was first detected in the amide N of glutamine, and only subsequently in the amino N of glutamate. Moreover, no assimilation was detected in the presence of the specific GS inhibitor methionine sulfoximine. These observations are consistent with a dominant role for

GS in the assimilation of ammonium in Sphagnum.

IT 56-65-5, 5'-ATP, biological studies

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(reaction with glutamine synthetase isoenzyme, kinetics of; ammonium assimilation in bryophytes: glutamine synthetase from Sphagnum fallax)

RN 56-65-5 CAPLUS

CN Adenosine 5'-(tetrahydrogen triphosphate) (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 6 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1996:323936 CAPLUS

DN 125:81261

TI Single-step signal group-imidazole labeling of organic phosphate groups under aqueous conditions

IN Giese, Roger W.; Wang, Poguang

PA Northeastern University, USA

SO U.S., 9 pp. CODEN: USXXAM

DT Patent

LA English

FAN. CNT 1

| FAN. | CNT 1 | | | | |
|------|------------------|------|----------|-----------------|----------|
| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
| | | | | | |
| PI | US 5512486 | Α | 19960430 | US 1993-60569 | 19930510 |
| OS | MARPAT 125:81261 | | | | |
| CT | | | | | |

Me
$$H_3$$
 CCONH

Me F F (CH₂) 2 CONHNHCOCHCH₂

NH

I

AB Compds. and methods for single-step, covalent labeling of the phosphate group of an org. substance under aq. conditions are described. The labeling compd. includes any kind of detectable signal group covalently bound to an **imidazole** moiety, which can be **imidazole** or a substituted **imidazole**. A preferred labeling compd. has the formula I.

IT 653-63-4 902-04-5 14490-86-9 148807-02-7 178388-84-6 RL: ANT (Analyte): RCT (Reactant):

RL: ANT (Analyte); RCT (Reactant); ANST (Analytical study); RACT (Reactant or reagent)

(1-step signal group-imidazole labeling of org. phosphate groups under aq. conditions)

653-63-4 CAPLUS RN

CN

5'-Adenylic acid, 2'-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

902-04-5 CAPLUS RN5'-Guanylic acid, 2'-deoxy- (9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

$$H_2N$$
 M
 H
 N
 N
 R
 R
 R
 OPO_3H_2
 OH

14490-86-9 CAPLUS RN5'-Guanylic acid, 8-(acetyl-9H-fluoren-2-ylamino)-2'-deoxy- (9CI) (CA CN INDEX NAME)

Absolute stereochemistry.

148807-02-7 CAPLUS RN5'-Adenylic acid, 2'-deoxy-7,8-dihydro-8-oxo- (9CI) (CA INDEX NAME)

RN 178388-84-6 CAPLUS

CN 5'-Adenylic acid, 2'-deoxy-8-(phenylmethoxy)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 7 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1992:585040 CAPLUS

DN 117:185040

TI Chemical modification and irreversible inhibition of striatal A2a adenosine receptors

AU Jacobson, Kenneth A.; Stiles, Gary L.; Ji, Xiao Duo

CS Lab. Bioorg. Chem., Natl. Inst. Diabetes Dig. Kidney Dis., Bethesda, MD, 20892, USA

SO Mol. Pharmacol. (1992), 42(1), 123-33 CODEN: MOPMA3; ISSN: 0026-895X

DT Journal

LA English

AB The ligand recognition site of A2a-adenosine receptors in rabbit striatal membranes was probed using non-site-directed labeling reagents and specific affinity labels. Exposure of membranes to di-Et pyrocarbonate at a concn. of 2.5 mM, followed by washing, was found to inhibit the binding of [3H]CGS 21680 and [3H]xanthine amine congener to A2a receptors, by 86 and 30%, resp. Protection from di-Et pyrocarbonate inactivation by an adenosine receptor agonist, 5'-N-ethylcarboxamidoadenosine, and an antagonist, theophylline, suggested the presence of two histidyl residues on the receptor, one assocd. with agonist binding and the other with antagonist binding. Binding of [3H]CGS 21680 or [3H]xanthine amine congener was partially restored after incubation with 250 mMhydroxylamine, further supporting histidine as the modification site. Preincubation with disulfide-reactive reagents, dithiothreitol or sodium dithionite, at >5 mM inhibited radioligand binding, indicating the presence of essential disulfide bridges in A2a receptors, whereas the concn. of mercaptoethanol required to inhibit binding was >50 mM. A no. of isothiocyanate-bearing affinity labels derived from the A2a-selective agonist 2-[(2-aminoethylamino)carbonylethylphenylethylamino]-5'-Nethylcarboxamidoadenosine (APEC) were synthesized and found to inhibit A2a receptor binding in rabbit and bovine striatal membranes. Binding to

rabbit al receptors was not inhibited. Preincubation with the affinity label 4-isothiocyanatophenylaminothiocarbonyl-APEC (100 nM) diminished the Bmax for [3H]CGS 21680 binding by 71%, and the Kd was unaffected, suggesting a direct modification of the ligand binding site. Reversal of 4-isothiocyanatophenylaminothiocarbonyl-APEC inhibition of [3H]CGS 21680 binding with hydroxylamine suggested that the site of modification by the isothiocyanate is a cysteine residue. A bromoacetyl deriv. of APEC was ineffective as an affinity label at submicromolar concns.

129666-43-9P 143999-46-6P 143999-47-7P TΤ 143999-48-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as purinergic P2a receptor affinity labeling probe)

129666-43-9 CAPLUS RN.beta.-D-Ribofuranuronamide, 1-[6-amino-2-[[2-[4-[3-[[2-[[(4-CNisothiocyanatophenyl)amino]thioxomethyl]amino]ethyl]amino]-3oxopropyl]phenyl]ethyl]amino]-9H-purin-9-yl]-1-deoxy-N-ethyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

143999-46-6 CAPLUS RN

.beta.-D-Ribofuranuronamide, 1-[6-amino-2-[[2-[4-[3-[[2-[[[3-CN

(aminocarbonyl) -5-isothiocyanatophenyl]amino]thioxomethyl]amino]ethyl]amino]-3-oxopropyl]phenyl]ethyl]amino]-9H-purin-9-yl]-1-deoxy-N-ethyl- (9CI)
(CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 143999-47-7 CAPLUS

CN

.beta.-D-Ribofuranuronamide, 1-[6-amino-2-[[2-[4-[3-[[2-[[[(3,5-disothiocyanatophenyl)amino]thioxomethyl]amino]ethyl]amino]-3-oxopropyl]phenyl]ethyl]amino]-9H-purin-9-yl]-1-deoxy-N-ethyl- (9CI) (CA INDEX NAME)

$$S = C = N$$

$$NH - C - NH - CH_2 - CH_2 - NH - C - CH_2 - CH_2$$

$$CH_2 - CH_2 -$$

PAGE 1-B

RN 143999-48-8 CAPLUS

.beta.-D-Ribofuranuronamide, 1-[6-amino-2-[[2-[4-[3-[[2-[[[(bromoacetyl)amino]thioxomethyl]amino]ethyl]amino]-3oxopropyl]phenyl]ethyl]amino]-9H-purin-9-yl]-1-deoxy-N-ethyl- (9CI) (CA INDEX NAME)

PAGE 1-A

IT 129681-42-1

RL: BIOL (Biological study)

(purinergic P2a receptor affinity labeling probe)

RN 129681-42-1 CAPLUS

CN .beta.-D-Ribofuranuronamide, 1-[6-amino-2-[[2-[4-[3-[[2-[[[(3-isothiocyanatophenyl)amino]thioxomethyl]amino]ethyl]amino]-3-oxopropyl]phenyl]ethyl]amino]-9H-purin-9-yl]-1-deoxy-N-ethyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

IT 126828-50-0

RL: RCT (Reactant)

(reaction of, with isothiocyanates, in prepn. of purinergic P2a receptor affinity labeling probes)

RN 126828-50-0 CAPLUS

CN .beta.-D-Ribofuranuronamide, 1-[6-amino-2-[[2-[4-[3-[(2-aminoethyl)amino]-3-oxopropyl]phenyl]ethyl]amino]-9H-purin-9-yl]-1-deoxy-N-ethyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

HO

 \gg_{OH}

L6 ANSWER 8 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1990:568692 CAPLUS

DN 113:168692

TI Conversion of 5-aminoimidazole ribotide to the pyrimidine of thiamin in enterobacteria: study of the pathway with specifically labeled samples of riboside

AU Estramareix, Bernard; David, Serge

CS Inst. Chim. Mol., Univ. Paris-Sud, Orsay, 91405, Fr.

SO Biochim. Biophys. Acta (1990), 1035(2), 154-60

CODEN: BBACAQ; ISSN: 0006-3002

DT Journal

LA English

AB Samples of 5-amino-1-(.beta.-D-ribofuranosyl)imidazole labeled with 13C at position C-1 or C-2 of the ribose part or with 15N at position N-3 or amino of the imidazole part were prepd. by chem. synthesis. The incorporation of label from these samples into the pyrimidine of thiamin biosynthesized by a mutant strain of Salmonella typhimurium was studied by GC-MS. The results show that in enterobacteria the Me carbon atom and the N-1 nitrogen atom of one mol. of thiamin pyrimidine derive from the same mol. of 5-aminoimidazole ribotide. More specifically, the Me carbon atom comes from the carbon C-2' of the ribose part and the nitrogen N-1 from nitrogen N-3 of the imidazole; furthermore, the amino nitrogen of the aminoimidazole becomes the amino

nitrogen of the pyrimidine.

25635-88-5 30597-39-8, 5-Aminoimidazole riboside IT

RL: PROC (Process)

(conversion of, to pyrimidine of thiamin by enterobacteria)

25635-88-5 CAPLUS RN

1H-Imidazol-5-amine, 1-(5-O-phosphono-.beta.-D-ribofuranosyl)- (9CI) (CA CNINDEX NAME)

Absolute stereochemistry.

30597-39-8 CAPLUS RN

1H-Imidazol-5-amine, 1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

129822-69-1P 129822-70-4P 129822-71-5P IT

129838-71-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

129822-69-1 CAPLUS RN

1H-Imidazole-4-carboxylic acid, 5-amino-1-(.beta.-D-ribofuranosyl-2-13C)-, CNethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

129822-70-4 CAPLUS RN

1H-Imidazole-3-15N-4-carboxylic acid, 5-amino-1-.beta.-D-ribofuranosyl-, CN (CA INDEX NAME) ethyl ester (9CI)

RN 129822-71-5 CAPLUS

CN 1H-Imidazole-4-carboxylic acid, 5-(amino-15N)-1-.beta.-D-ribofuranosyl-,
 methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HO OH
$$15_{\mathrm{NH}_2}$$
 OMe

RN 129838-71-7 CAPLUS

CN 1H-Imidazole-4-carboxylic acid, 5-amino-1-(.beta.-D-ribofuranosyl-1-13C)-, ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 9 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1988:438155 CAPLUS

DN 109:38155

TI Synthesis of fluorescent or biotinylated nucleoside compounds

AU Sarfati, S. R.; Pochet, Sylvie; Guerreiro, C.; Namane, A.; Huynh Dinh, Tam; Igolen, Jean

CS Dep. Biochim. Genet. Mol., Inst. Pasteur, Paris, 75724/15, Fr.

SO Tetrahedron (1987), 43(15), 3491-7 CODEN: TETRAB; ISSN: 0040-4020

DT Journal

LA English

OS CASREACT 109:38155

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Two types of modified nucleoside derivs. I (R = H, dansyl) of 2'-deoxycytidine and II of 2'-deoxyadenosine, useful for the specific attachment of non-radioactive labeling reagents such as fluorescent or

biotinyl group were prepd. II converted into the biotinylated deriv. which is a substrate for DNA polymerase.

IT 115244-09-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and benzoylation of)

RN 115244-09-2 CAPLUS

CN Carbamic acid, [10-[[6-amino-9-(2-deoxy-.beta.-D-erythro-pentofuranosyl)-9H-purin-8-yl]amino]decyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 115244-11-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and conversion of, to morpholide)

RN 115244-11-6 CAPLUS

CN Carbamic acid, [10-[[6-amino-9-(2-deoxy-5-0-phosphono-.beta.-D-erythro-pentofuranosyl)-9H-purin-8-yl]amino]decyl]-, C-(phenylmethyl) ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 115244-16-1P

RN 115244-16-1 CAPLUS

CN Carbamic acid, [10-[[6-(benzoylamino)-9-(3,5-di-0-benzoyl-2-deoxy-.beta.-D-erythro-pentofuranosyl)-9H-purin-8-yl]amino]decyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

IT 115244-14-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and hydrogenolysis of)

RN 115244-14-9 CAPLUS

CN Carbamic acid, [10-[[6-amino-9-[2-deoxy-5-0-(hydroxy-4-morpholinylphosphinyl)-.beta.-D-erythro-pentofuranosyl]-9H-purin-8-yl]amino]decyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 115244-15-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and phosphorylation of)

RN 115244-15-0 CAPLUS

CN Adenosine, 8-[(10-aminodecyl)amino]-2'-deoxy-, 5'-(hydrogen 4-morpholinylphosphonate) (9CI) (CA INDEX NAME)

IT 115244-12-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with biotin deriv.)

RN 115244-12-7 CAPLUS

CN Adenosine 5'-(tetrahydrogen triphosphate), 8-[(10-aminodecyl)amino]-2'-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 115260-06-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with cyanoethyl phosphate)

RN 115260-06-5 CAPLUS

CN Carbamic acid, [10-[[6-(benzoylamino)-9-(3-0-benzoyl-2-deoxy-.beta.-D-erythro-pentofuranosyl)-9H-purin-8-yl]amino]decyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 115244-08-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with imidazole deriv.)

RN 115244-08-1 CAPLUS

CN Adenosine, 8-[(10-aminodecyl)amino]-2'-deoxy- (9CI) (CA INDEX NAME)

IT 115244-10-5P

RN 115244-10-5 CAPLUS

CN Carbamic acid, [10-[[6-(benzoylamino)-9-(2-deoxy-.beta.-D-erythro-pentofuranosyl)-9H-purin-8-yl]amino]decyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 115244-13-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as nonradioactively labeled deoxyadenosine deriv.)

RN 115244-13-8 CAPLUS

CN Adenosine 5'-(tetrahydrogen triphosphate), 2'-deoxy-8-[[10-[[5-(hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl)-1-oxopentyl]amino]decyl]amino]-, [3aS-(3a.alpha.,4.beta.,6a.alpha.)]- (9CI) (CA INDEX NAME)

IT 14985-44-5

RL: RCT (Reactant)

(reaction of, with decanediamine)

RN14985-44-5 CAPLUS

Adenosine, 8-bromo-2'-deoxy- (7CI, 8CI, 9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

L6 ANSWER 10 OF 23 CAPLUS COPYRIGHT 2002 ACS

1987:571439 CAPLUS ΑN

DN 107:171439

ΤI Studies of the functional topography of Escherichia coli RNA polymerase. Affinity labelling of RNA polymerase in a promoter complex by phosphorylating derivatives of primer oligonucleotides

ΑU Godovikova, T. S.; Grachev, M. A.; Kutyavin, I. V.; Tsarev, I. G.; Zarytova, V. F.; Zaychikov, E. F.

CS

Novosibirsk Inst. Bioorg. Chem., Novisibirsk, 630090, USSR Eur. J. Biochem. (1987), 166(3), 611-16 SO CODEN: EJBCAI; ISSN: 0014-2956

DTJournal

LAEnglish

AB Amidation of the 5'-phosphate group of the heptanucleotide pdApdApdApdTpdCpdGprC and of its derivs. of the general formula (pdN) npdGprC (n = 0-5, dN = deoxynucleoside) with imidazole,N-methylimidazole, and 4-dimethylaminopyridine afforded a series of phosphorylating affinity reagents. The parent oligonucleotides of this series are complementary to promoter A2 of T7 phage over the region (-5 to +2) and are known to be efficient primers of the synthesis of RNA by E. coli RNA polymerase with promoter A2 as template. Treatment of the

complex RNA-polymerase.cntdot.promter-A2 with affinity reagents followed by addn. of [.alpha.-32P]UTP resulted in labeling of RNA polymerase by the residues -(pdN)npdGprC*prU (*p = radioactive phosphate). This affinity labeling was highly selective because elongation of the covalently bound residues (pdN)npdGprC by *prU residues was catalyzed by the active center of RNA polymerase. The most efficient reagents were N-methylimidazolides. A dramatic change of the pattern of labeling of the subunits .beta., .beta.', and .sigma. took place with changing n. Max. labeling of the .beta. subunit occurred at n = 1 and of the .sigma. subunit at n = 5. The targets in both the subunits were histidine residues. The .alpha. subunit was not specifically labeled.

IT 94479-02-4 94479-07-9 94479-08-0 94479-09-1 110651-95-1 110671-51-7

RL: RCT (Reactant)
(amidation of)

RN 94479-02-4 CAPLUS

CN

Cytidine, 2'-deoxy-5'-O-phosphonocytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

94479-07-9 CAPLUS

RN

CN

Cytidine, 5'-O-phosphonothymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

RN 94479-08-0 CAPLUS

CN Cytidine, 2'-deoxy-5'-O-phosphonoadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

RN 94479-09-1 CAPLUS

CN Cytidine, 2'-deoxy-5'-0-phosphonoadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c|c}
OH & & & \\
\hline
OPO & & & \\
0 & & & \\
H_2O_3PO-CH_2 & & & \\
NH_2
\end{array}$$

$$\begin{array}{c|c} & & \\ & &$$

RN 110651-95-1 CAPLUS

CN Cytidine, 2'-deoxy-5'-O-phosphonoguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$H_2N$$
 H_2N
 H_2N

RN 110671-51-7 CAPLUS

CN Cytidine, 2'-deoxy-5'-O-phosphonoadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 2-B

PAGE 1-B

__Me

PAGE 2-A

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 107759-25-1 CAPLUS

CN Cytidine, 2'-deoxy-5'-O-[hydroxy(3-methyl-1H-imidazolium-1-yl)phosphinyl]adenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-, inner salt (9CI) (CA INDEX NAME)

PAGE 1-A

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 107759-26-2 CAPLUS

CN Adenosine, cytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy-, 5'-[hydrogen (3-methyl-1H-imidazolium-1-yl)phosphonate], inner salt (9CI) (CA INDEX NAME)

PAGE 2-B

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 107787-13-3 CAPLUS

CN Cytidine, 5'-O-[hydroxy(3-methyl-1H-imidazolium-1-yl)phosphinyl]guanylyl-(3'.fwdarw.5')-, inner salt (9CI) (CA INDEX NAME)

$$\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

PAGE 2-A

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 107787-14-4 CAPLUS

CN Cytidine, 2'-deoxy-5'-O-[hydroxy(3-methyl-1H-imidazolium-1-yl)phosphinyl]cytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-, inner salt (9CI) (CA INDEX NAME)

PAGE 2-A

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 107787-15-5 CAPLUS

Adenosine, cytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxy-, 5'-[hydrogen (3-methyl-1H-imidazolium-1-yl)phosphonate], inner salt (9CI) (CA INDEX NAME)

*** FRAGMENT DIAGRAM IS INCOMPLETE ***

RN 110651-96-2 CAPLUS

CN Cytidine, 2'-deoxy-5'-O-(hydroxy-1H-imidazol-1-ylphosphinyl)guanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

PAGE 1-A

PAGE 2-A

110651-98-4 CAPLUS RN

Cytidine, 5'-0-(hydroxy-1H-imidazol-1-ylphosphinyl)thymidylyl-CN(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5') - (9CI) (CA INDEX NAME)

RN 110651-99-5 CAPLUS
CN Cytidine, 2'-deoxy-5'-O-(hydroxy-1H-imidazol-1-ylphosphinyl)adenylyl(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

$$N = 0$$

$$N =$$

RN 110652-00-1 CAPLUS
CN Cytidine, 2'-deoxy-5'-O-(hydroxy-1H-imidazol-1-ylphosphinyl)adenylyl(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI)
(CA INDEX NAME)

PAGE 1-A

RN 110652-01-2 CAPLUS
CN Cytidine, 2'-deoxy-5'-O-(hydroxy-1H-imidazol-1-ylphosphinyl)adenylyl(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 2-B

RNCN

110652-02-3 CAPLUS Cytidine, 2'-deoxy-5'-O-[(dimethylamino)hydroxyphosphinyl]guanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 2-A

110652-03-4 CAPLUS RNCN

Cytidine, 2'-deoxy-5'-O-[(dimethylamino)hydroxyphosphinyl]adenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

RN 110652-04-5 CAPLUS
CN Cytidine, 2'-deoxy-5'-O-[(dimethylamino)hydroxyphosphinyl]adenylyl(3'.fwdarw.5')-2'-deoxyadenylyl-(3'.fwdarw.5')-2'-deoxyadenylyl(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-B

RN

CN

110671-52-8 CAPLUS
Cytidine, 2'-deoxy-5'-O-[(dimethylamino)hydroxyphosphinyl]cytidylyl(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

PAGE 2-A

RN 110671-53-9 CAPLUS
CN Cytidine, 5'-O-[(dimethylamino)hydroxyphosphinyl]thymidylyl-(3'.fwdarw.5')2'-deoxycytidylyl-(3'.fwdarw.5')-2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI)
(CA INDEX NAME)

PAGE 2-A

$$\begin{array}{c} NH_2 \\ NH_2 \\ NH_2 \\ NH_3 \\ NH_4 \\ NH_5 \\ NH_6 \\ NH$$

RN 110671-54-0 CAPLUS
CN Cytidine, 2'-deoxy-5'-O-[(dimethylamino)hydroxyphosphinyl]adenylyl(3'.fwdarw.5')-thymidylyl-(3'.fwdarw.5')-2'-deoxycytidylyl-(3'.fwdarw.5')2'-deoxyguanylyl-(3'.fwdarw.5')- (9CI) (CA INDEX NAME)

L6 ANSWER 11 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1984:420330 CAPLUS

DN 101:20330

TI Biosynthesis of thiamin: 5-aminoimidazole ribotide as the precursor of all the carbon atoms of the pyrimidine moiety

AU Estramareix, Bernard; Therisod, Michel

CS Lab. Chim. Org. Multifonct., Univ. Paris-Sud, Orsay, 91405, Fr.

SO J. Am. Chem. Soc. (1984), 106(13), 3857-60 CODEN: JACSAT; ISSN: 0002-7863

DT Journal

LA English

Three C atoms of the pyrimidine moiety of thiamin originate in the AΒ imidazole ring of 5-aminoimidazole ribotide (AIR) in bacteria. The origin of the other 3 C atoms was examd. with labeled biosynthetic samples of 5-aminoimidazole riboside (AIRs). The incorporation of label was studied in a Salmonella typhimurium strain able to synthesize the pyrimidine part of thiamin from glucose and a minute amt. of AIRs. No incorporation of 13C was found in the pyrimidine synthesized from [U-13C] glucose and natural AIRs. In contrast, the isotopic compn. of the pyrimidine synthesized from natural glucose and [U-13C]AIRs was close to that of the labeled AIRs. From AIRs labeled mainly in its ribose part with 14C and inactive glucose, a pyrimidine labeled mainly in the 3 C atoms that do not derive from the imidazole part was obtained. These C atoms were approx. as radioactive as those of the ribose part of AIRs. It was concluded that the 3 C atoms of the pyrimidine moiety of thiamin originate in the ribose part of AIRs, which is, thus, the precursor of all the C atoms of this pyrimidine.

IT 25635-88-5

RL: BIOL (Biological study)

(thiamin pyrimidine moiety carbon atoms derived from)

RN 25635-88-5 CAPLUS

CN 1H-Imidazol-5-amine, 1-(5-O-phosphono-.beta.-D-ribofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 12 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1983:194610 CAPLUS

DN 98:194610

TI Quantitative determination of adenosine

IN Sato, Tomokazu; Ui, Michio

PA Yamasa Shoyu Co., Ltd., Japan

SO Eur. Pat. Appl., 24 pp. CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

| | PATENT NO. | | KIND | DATE | APF | LICATION N | 10. | DATE |
|------|------------|---------|---------|----------|-----|------------|-----|----------|
| | | | | | | | | |
| ΡI | EP 70033 | 1 | A1 | 19830119 | EP | 1982-10627 | 6 | 19820713 |
| | EP 70033 | | B1 | 19841121 | | | | |
| | R: | CH, DE, | FR, GB, | LI | | | | |
| | JP 58011 | .857 | A2 | 19830122 | JP | 1981-11132 | 4 | 19810715 |
| | JP 62057 | 220 | B4 | 19871130 | | | | |
| | US 44789 | 34 | Α | 19841023 | US | 1982-39686 | 3 | 19820709 |
| | CA 11777 | '50 | A1 | 19841113 | CA | 1982-40725 | 4 | 19820714 |
| PRAI | JP 1981- | 111324 | | 19810715 | | | | |

An accurate and sensitive immunoassay is described for the detn. of adenosine in biol. fluids following its conversion to 2',3'-diacyladenosine (by an acid anhydride in the presence of an org. tert-amine) by using antibodies specific for 2',3'-diacyladenosine and labeled 2',3'-diacyladenosine. The antibodies were prepd. by inoculation of an animal with an antigen consisting of the condensation product of 2',3'-diacyladenosine and a carrier protein (e.g. human serum albumin) via dicarboxylic acid residues. Thus, adenosine was detd. in rat plasma treated with MnCl2 and benzyl alc. by a RIA. An acetylating agent consisting of succinic anhydride and triethylamine was added to the sample and stds., followed by diln. with imidazole buffer (pH 5-8), mixing with a predetd. amt. of 2',3'-disuccinyladenosine-3H and prepd. antibodies (sol. or immobilized), sepn. of bound and free label with dextran-coated carbon, and radioactivity detn.

IT 58-61-7, analysis

RL: ANT (Analyte); ANST (Analytical study)
 (detn. of, in biol. fluids by immunoassay)

RN 58-61-7 CAPLUS

CN Adenosine (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 84872-85-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, for adenosine detn. in body fluids by immunoassay)

RN 84872-85-5 CAPLUS

CN Adenosine, 2',3'-bis(hydrogen butanedioate) (9CI) (CA INDEX NAME)

L6 ANSWER 13 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1983:122427 CAPLUS

DN 98:122427

TI Stabilization of glucose oxidase apoenzyme

IN Rupchock, Patricia A.; Tyhach, Richard J.

PA Miles Laboratories, Inc., USA

SO U.S., 17 pp. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

11S 4366243 A 19821228 US 1981-255310 19810417

US 1981-255310 19821228 PΙ US 4366243 Α Glucose oxidase apoenzyme is stabilized by poly(vinyl alc.) and serum AB albumin for ligand binding assays. The stabilized apoenzyme can be incorporated into test strips for immunoassays. In such assays an FAD-antigen conjugate is the label, and FAD-antigen conjugate which is not bound to the antibody is available for glucose oxidase apoenzyme activation. For example, test strips were prepd. for dinitrophenyl caproate immunoassay which contained buffer, a glucose oxidase detection system, apoglucose oxidase, dinitrophenol antibody, and dinitrophenol-FAD conjugate. Inclusion of poly(vinyl alc.) and albumin increased the heat stability of the test strips. Test strips for theophylline and phenytoin are also described.

IT 76748-73-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with dinitrofluorobenzene)

RN 76748-73-7 CAPLUS

CN Riboflavin 5'-(trihydrogen diphosphate), P'.fwdarw.5'-ester with N-(6-aminohexyl)adenosine (9CI) (CA INDEX NAME)

IT 73122-00-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with riboflavin monophosphate)

RN 73122-00-6 CAPLUS

CN Adenosine, N-[6-[(trifluoroacetyl)amino]hexyl]-, 5'-(hydrogen 1H-imidazol-1-ylphosphonate) (9CI) (CA INDEX NAME)

IT 82604-51-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as label for amino assay with apoglucose oxidase)

RN 82604-51-1 CAPLUS

CN 5'-Adenylic acid, N-[6-[(2,4-dinitrophenyl)amino]hexyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 146-14-5DP, reaction products with antigens

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as **label** for immunoassay with stabilized apoglucose oxidase)

RN 146-14-5 CAPLUS

CN Riboflavin 5'-(trihydrogen diphosphate), P'.fwdarw.5'-ester with adenosine (9CI) (CA INDEX NAME)

IT 66060-76-2

RL: RCT (Reactant)

(reaction of, with carbonyldiimidazole)

RN 66060-76-2 CAPLUS

CN 5'-Adenylic acid, N-[6-[(trifluoroacetyl)amino]hexyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 14 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1982:452162 CAPLUS

DN 97:52162

TI Homogeneous specific binding assay device and an analytical method using the device

IN Greenquist, Alfred C.; Li, Thomas M.; Rupchock, Patricia A.; Tyhach,
Richard Joseph; Walter, Bert

PA Miles Laboratories, Inc., USA

SO Eur. Pat. Appl., 93 pp. CODEN: EPXXDW

DT Patent

LA English

FAN. CNT 1

| FAN. | CNT 1 | | | | | | | |
|------|-------|-------|-----|--------|-----------|-----|-----------------|----------|
| | PATEN | T NO. | | KIND | DATE | | APPLICATION NO. | DATE |
| DT | BD 61 | 212 | | | | | | |
| ΡI | EP 51 | 213 | | A1 | 19820512 | | EP 1981-108681 | 19811022 |
| | EP 51 | 213 | | B1 | 19870325 | | | |
| | R | : AT, | ΒE, | CH, DE | , FR, GB, | IT, | LU, NL, SE | |
| | CA 11 | 83450 | | A1 | 19850305 | | CA 1981-381675 | 19810714 |
| | IL 63 | 333 | | A1 | 19860131 | | IL 1981-63333 | 19810716 |
| | IL 74 | 590 | | A1 | 19860131 | | IL 1981-74590 | 19810716 |
| | IL 74 | 591 | | A1 | 19860131 | | IL 1981-74591 | 19810716 |
| | ZA 81 | 04949 | | A | 19820929 | | ZA 1981-4949 | 19810720 |

| | AU | 8175047 | A1 | 19820610 | ΑU | 1981-75047 | 19810908 |
|------|----|-------------|-----|----------|----|-------------|----------|
| | ΑU | 530707 | B2 | 19830728 | | | |
| | NO | 8103483 | Α | 19820503 | NO | 1981-3483 | 19811015 |
| | NO | 161703 | В | 19890605 | | | |
| | NO | 161703 | C | 19890913 | | | |
| | ΑT | 26187 | E | 19870415 | AΤ | 1981-108681 | 19811022 |
| | FI | 8103369 | Α | 19820501 | FI | 1981-3369 | 19811028 |
| | FI | 75677 | В | 19880331 | | | |
| | FI | 75677 | C | 19880711 | | | |
| | DK | 8104784 | Α . | 19820501 | DK | 1981-4784 | 19811029 |
| | DK | 157328 | В | 19891211 | | | |
| | DK | 157328 | С | 19900507 | | | |
| | JΡ | 57103055 | A2 | 19820626 | JP | 1981-172169 | 19811029 |
| | JP | 02035261 | B4 | 19900809 | | | |
| | ES | 506689 | A1 | 19830401 | ES | 1981-506689 | 19811029 |
| | ES | 518735 | A1 | 19840201 | ES | 1982-518735 | 19821231 |
| | ES | 518736 | A1 | 19840201 | ES | 1982-518736 | 19821231 |
| | ES | 518737 | A1 | 19840201 | ES | 1982-518737 | 19821231 |
| | ES | 518738 | A1 | 19840201 | ES | 1982-518738 | 19821231 |
| | ES | 518739 | A1 | 19840201 | ES | 1982-518739 | 19821231 |
| | | 4668619 | Α | 19870526 | US | 1984-632946 | 19840720 |
| PRAI | US | 1980-202378 | | 19801030 | | | |
| | | 1981-63333 | | 19810716 | | | |
| | | 1981-108681 | | 19811022 | | | |
| | US | 1982-381218 | | 19820524 | | | |
| | | | | | | | |

AB A binding assay test element is described (e.g., for antibody or antigen immunoassays) which has a solid carrier (e.g., paper, polymeric film, or gel) impregnated with reagents. The assay systems include those involving enzyme substrate labels, enzyme prosthetic group labels, and enzyme labels. A detectable response (e.g., luminescent, fluorescent, spectrophotometric, or colorimetric) is producted which is a function of the amt. of analyte. Examples include theophylline detn. with theophylline-FAD conjugate as label and glucose oxidase apoenzyme, galactosyl-umbelliferone-theophylline conjugate as label and .beta.-galactosidase, and glucose 6-phosphate dehydrogenase-theophylline conjugate as label. Further examples include detns. of N-(2,4-dinitrophenyl)-.epsilon.-aminocaproic acid and other drugs.

IT 73122-00-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with riboflavin monophosphate)

RN 73122-00-6 CAPLUS

CN Adenosine, N-[6-[(trifluoroacetyl)amino]hexyl]-, 5'-(hydrogen lH-imidazol-1-ylphosphonate) (9CI) (CA INDEX NAME)

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of, as label for dinitrophenylaminocaproic acid
 immunoassay)

RN 82604-51-1 CAPLUS

CN 5'-Adenylic acid, N-[6-[(2,4-dinitrophenyl)amino]hexyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 76748-73-7 CAPLUS

CN Riboflavin 5'-(trihydrogen diphosphate), P'.fwdarw.5'-ester with N-(6-aminohexyl)adenosine (9CI) (CA INDEX NAME)

PAGE 1-A

IT 66060-76-2

RL: RCT (Reactant)

(reaction of, with carbonyldiimidazole)

RN 66060-76-2 CAPLUS

CN 5'-Adenylic acid, N-[6-[(trifluoroacetyl)amino]hexyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L6 ANSWER 15 OF 23 CAPLUS COPYRIGHT 2002 ACS
- AN 1982:196189 CAPLUS
- DN 96:196189
- TI Homogeneous specific binding assay employing an intramolecularly modulated photogenic enzyme substrate **label**
- IN Burd, John F.; Li, Thomas M.
- PA Miles Laboratories, Inc., USA
- SO U.S., 14 pp. CODEN: USXXAM
- DT Patent
- LA English
- FAN.CNT 1

PATENT NO. KIND DATE

APPLICATION NO. DATE

- PI US 4318981 A 19820309 US 1980-143497 19800424
- AB Binding assays (e.g., immunoassays) are described that use a labeled conjugate which is cleavable by an enzyme to produce an indicator which emits light. The label component of the labeled conjugate consists of a photophore, a linkage which is cleavable by an enzyme, a modulator for light emission by the photophore, and a linking group through which the label component is bound to the rest of the conjugate. An example is given of fluorescence immunoassay of theophylline with FAD-theophylline as labeled conjugate and nucleotide pyrophosphatase as enzyme. The flavin portion of FAD serves as fluorescer, the adenosine moiety of FAD serves as quencher, and the pyrophosphate linking group in FAD serves as enzyme-cleavable linkage.

IT 39007-51-7

RL: ANST (Analytical study)
 (as fluorescing agent, in fluorescence binding assay with enzyme
 substrate labels)

RN 39007-51-7 CAPLUS

CN 3H-Imidazo[2,1-i]purine, 3-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 76748-73-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with dimethyltetrahydropyridopurinetrione)

RN 76748-73-7 CAPLUS

CN Riboflavin 5'-(trihydrogen diphosphate), P'.fwdarw.5'-ester with N-(6-aminohexyl)adenosine (9CI) (CA INDEX NAME)

PAGE 1-A

IT 73122-00-6P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with riboflavin monophosphate)

RN

73122-00-6 CAPLUS
Adenosine, N-[6-[(trifluoroacetyl)amino]hexyl]-, 5'-(hydrogen CN1H-imidazol-1-ylphosphonate) (9CI) (CA INDEX NAME)

IT 146-14-5DP, reaction products with theophylline RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, for fluorescence immunoassay)

RN 146-14-5 CAPLUS

CN Riboflavin 5'-(trihydrogen diphosphate), P'.fwdarw.5'-ester with adenosine (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 66060-76-2

RL: RCT (Reactant)

(reaction of, with carbonyl diamidazole)

RN 66060-76-2 CAPLUS

CN 5'-Adenylic acid, N-[6-[(trifluoroacetyl)amino]hexyl]- (9CI) (CA INDEX NAME)

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L6 ANSWER 16 OF 23 CAPLUS COPYRIGHT 2002 ACS
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AN 1981:116179 CAPLUS

DN 94:116179

TI Spin-labeling study of intermolecular interactions and self-organization of nucleotide systems into ordered structures

AU Petrov, A. I.; Sukhorukov, B. I.

CS Inst. Biol. Phys., Moscow, USSR

SO Stud. Biophys. (1980), 80(2), 79-84 CODEN: STBIBN; ISSN: 0081-6337

DT Journal

LA English

AB With the use of the spin label N-(2,2,5,5-tetramethyl-3-carbonylpyrroline-1-oxyl) imidazole it was demonstrated that: (1) the self assocn. of adenine nucleotides (AMP, ADP, or ATP) depended on protonation state and not on phosphate chain length; (2) protonation-induced changes in poly(A), poly(U), or poly(C) occurred in an alternating fashion, i.e., if the initial rigid structure of the polynucleotide changed to another rigid structure, there was an intermediate protonation degree at which the polynucleotide existed in a flexible conformation; and (3) complex formation between adenosine, uridine, or cytidine and poly(U) occurred with a free energy of H-bonding of 1.77, 0.86, and 0.51 kcal/mol, resp.

IT 24937-83-5

RL: BIOL (Biological study)
 (mononucleotide interaction with)

RN 24937-83-5 CAPLUS

CN 5'-Adenylic acid, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 61-19-8 CMF C10 H14 N5 O7 P

Absolute stereochemistry.

RN 58-61-7 CAPLUS

CN Adenosine (8CI, 9CI) (CA INDEX NAME)

IT 56-65-5, biological studies 58-64-0, biological studies

61-19-8, biological studies

RL: PRP (Properties)

(self assocn. of, phosphate and protonation effect on)

RN 56-65-5 CAPLUS

CN Adenosine 5'-(tetrahydrogen triphosphate) (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 58-64-0 CAPLUS

CN Adenosine 5'-(trihydrogen diphosphate) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 61-19-8 CAPLUS

CN 5'-Adenylic acid (8CI, 9CI) (CA INDEX NAME)

L6 ANSWER 17 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1981:11745 CAPLUS

DN 94:11745

TI Spin-labeled polyribonucleotides

AU Petrov, A. I.; Sukhorukov, B. I.

CS Inst. Biol. Phys., Pushchino, 142292, USSR

SO Nucleic Acids Res. (1980), 8(18), 4221-34 CODEN: NARHAD; ISSN: 0305-1048

DT Journal

LA English

AB Poly(U), poly(C), and poly(A) were spin-labeled with N-(2,2,5,5-tetramethyl-3-carbonylpyrroline-1-oxyl)imidazole. This spin label interacts selectively with the 2'-OH of ribose groups of polynucleotides and does not modify the nucleic acid bases. The extent of spin-labeling is not dependent on the nature of the base and is entirely detd. by rigidity of the secondary structure of the polynucleotide. The extent of modification for poly(U), poly(C), and poly(A) was 4.2, 1.7, and 1.5%, resp., the secondary structure of the polynucleotides being practically unchanged. Some physicochem. properties of the spin-labeled polynucleotides were investigated by ESR spectroscopy. Rotational correlation times of the spin label and activation energy of its motion were calcd.

IT 24937-83-5

RL: BIOL (Biological study)

(spin labeling of ribose of, ESR in relation to)

RN 24937-83-5 CAPLUS

CN 5'-Adenylic acid, homopolymer (9CI) (CA INDEX NAME)

CM 1

CRN 61-19-8

CMF C10 H14 N5 O7 P

Absolute stereochemistry.

L6 ANSWER 18 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1979:400478 CAPLUS

DN 91:478

TI In vitro effects of ionophore A23187 on skeletal collagen and noncollagen protein synthesis

AU Dietrich, John W.; Paddock, David N.

CS Sch. Med., Univ. Illinois, Peoria, IL, 61605, USA

SO Endocrinology (Baltimore) (1979), 104(2), 493-9 CODEN: ENDOAO; ISSN: 0013-7227

DT Journal

LA English

GΙ

AΒ Effects of ionophore A 23187 (I) [52665-69-7] on skeletal collagen formation were investigated in vitro. Collagen synthesis was quantitated in fetal rat calvaria by measuring proline-3H incorporation into collagenase-digestible (CDP) and noncollagen protein (NCP) using purified bacterial collagenase. I (0.03-1.0 .mu.g/mL) inhibited incorporation of label into CDP and NCP after 24 h of culture, with a greater effect on CDP. The response was not assocd. with altered amino acid uptake, precursor pool size, or degrdn. of newly labeled protein. Submaximal concns. of I and parathyroid hormone [9002-64-6] or dibutyryl [362-74-3] decreased CDP formation to a greater extent than treatment with the agents alone. Imidazole [288-32-4] although ineffective by itself, enhanced the effect of I. Alteration of medium Ca did not affect the response to I. The inhibitory effect of I was partially reversed by 24 h and completely reversed by 48 h of control treatment subsequent to an initial 24-h incubation with I. Indomethacin had no effect on CDP or NCP formation, either in the presence or absence of I. I did not alter the uptake of thymidine-3H or uridine-3H into acid-extractable pools but decreased incorporation of label into DNA and RNA, resp. Histol. examn. showed no difference between control and I treatment after 24 h. Apparently, I decreases bone collagen and noncollagen protein synthesis, possibly through a Ca-mediated effect. The mechanism of the inhibitory effect on DNA and RNA labeling is unknown, although it may be related to Ca. Ca may be involved in the actions of parathyroid hormone and dibutyryl cAMP on skeletal collagen synthesis.

I

IT 362-74-3

RL: BIOL (Biological study)

(collagen and protein formation response to, in bone)

RN 362-74-3 CAPLUS

CN Adenosine, N-(1-oxobutyl)-, cyclic 3',5'-(hydrogen phosphate) 2'-butanoate (9CI) (CA INDEX NAME)

L6 ANSWER 19 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1979:134457 CAPLUS

DN 90:134457

TI Cobalt(III) labeling of methionyl-tRNA synthetase from Escherichia coli

AU Kalogerakos, Theodore; Blanquet, Sylvain; Waller, Jean Pierre

CS Lab. Biochim., Ec. Polytech., Palaiseau, Fr.

SO Eur. J. Biochem. (1979), 93(2), 339-43

CODEN: EJBCAI; ISSN: 0014-2956

DT Journal

LA English

AB Native and trypsin-modified methionyl-tRNA synthetases from E. coli were inactivated by incubation in the presence of Co(III) complexes of ATP, stabilized either by imidazole or phenanthroline, or by oxidn. in situ to Co(III) of the substrate, ATP-Co(II). The inactivation proceeded by specific labeling of the catalytic ATP-Mg(II) site of the synthetases. The enzymes were completely inactivated by the incorporation of 1 Co and 1 ATP/active site. The inactivated enzymes were stored for a long period without significant reactivation or removal of the Co label. In the presence of dithiothreitol or 2-mercaptoethanol, the labeled enzymes recovered full activity with concomittant release of the bound label mols.

IT 56-65-5D, cobalt(III) complexes

RL: BIOL (Biological study)

(methionyl-tRNA synthetase inactivation by)

RN 56-65-5 CAPLUS

CN Adenosine 5'-(tetrahydrogen triphosphate) (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1973:132883 CAPLUS

DN 78:132883

TI Relaxation spectra of aspartate transcarbamylase. Interaction of the native enzyme with an adenosine 5'-triphosphate analog

Wu, Cheng-Wen; Hammes, Gordon G. ΔII

Dep. Chem., Cornell Univ., Ithaca, N. Y., USA

CS Biochemistry (1973), 12(7), 1400-8 SO

CODEN: BICHAW

Journal DT

English LA

AB

The interaction of aspartate transcarbamylase from Escherichia coli with the activator 6-mercapto-9-.beta.-D-ribofuranosylpurine 5'-triphosphate (sRTP) was studied at pH 7.0, 25.degree., in 0.15.MU. KOaac-0.04.MU. imidazole acetate, using difference spectroscopy and the temp.-jump method. The sRTP does not serve as an affinity label for the regulatory or catalytic sites, but a difference spectrum is obsd. when sRTP binds to the catalytic subunit, the regulatory subunit and the native enzyme. A spectral titration of the catalytic subunit indicates 3 binding sites are present per catalytic subunit mol. with a dissociation constant of 2.5 .times. 10-4m. With native enzyme, 2 relaxation processes are seen. The faster one has a time constant similar to that found with the isolated catalytic subunit and disappears in the presence of 2m.MU. carbamyl phosphate so that it probably reflects the interaction of sRTP catalytic site. The reciprocal relaxation time for the slower process increases and approaches a constant value as the sRTP concn. is raised. This behavior is obsd. in the presence or absence of 2m.MU. carbamyl phosphate and 10m.MU. succinate, although the limiting value reached varies. The simplest mechanism consistent with the data is a rapid combination of sRTP and enzyme followed by a rate-limiting conformational change, a mechanism similar to that proposed for the interaction of CTP with the native enzyme. When both sRTP and 5-bromocytidine 5-triphosphate are added to the enzyme, only a single relaxation process is obsd., suggesting that the same 2 conformational states occur with both activator and inhibitor complexes. A multiconformational model involving both concerted and sequential conformation transitions is proposed for the overall regulatory mechanism.

IT 27652-34-2

RL: PROC (Process)

(aspartate transcarbamylase binding of)

27652-34-2 CAPLUS RN

Inosine 5'-(tetrahydrogen triphosphate), 6-thio- (9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

ANSWER 21 OF 23 CAPLUS COPYRIGHT 2002 ACS L6

1966:13527 CAPLUS AN

64:13527 DN

OREF 64:2512h,2513a-b

Evidence for conformation changes in actin on contraction TТ

Szent-Gyorgyi, Andrew G. ΑU

Dartmouth Med. School, Hanover, NH CS

Muscle, Proc. Symp., Edmonton, Alberta, Can. (1965), 1964, 141-51 SO

DTJournal

LΑ English

Actomyosin (I), prepd. from F actin (II) contg. ADP labeled with 32P, AB suspended in pH 7.0 imidazole-HCl buffer contg. NaCl and MgCl2, was mixed with ATP. Aliquots were removed at intervals, centrifuged, and supernatants assayed for radioactivity, inorg. phosphate, and protein. Up to 25% total label was released in 30 sec., accompanied by ATP hydrolysis and superpptn. of protein. A further 25% of label was released at a much slower rate. Release of label required ATP. No reabsorption of label after ATP exhaustion was noted. II labeled with ATP-14C was incubated with 100-fold excess of ATP or an ATP-phosphocreatine (III) -creatine kinase mixt. in NaCl concns. of 0.05, 0.1, and 0.15M for 30-60 min., centrifuged for 3 hrs. at 40,000 rpm., and radioactivity and protein detd. in the supernatant. About 20% nucleotides and 8-15% protein did not sediment. I prepd. from labeled II retained its sp. activity after repeated repptns. Labeled ADP introduced with II into I was protected from creatine kinase action. Release of nucleotide from I was not catalyzed by ADP or AMP. At 10.degree., as opposed to 23.degree., superpptn. of I and release of nucleotide was delayed 10 min. Addn. of III enhanced the temp.-dependent effect. Creatine release was 10 times faster at 23.degree. than at 10.degree.. Superpptn., rate of release of nucleotide, and ATPase activity were all accelerated by Mg++. The data suggest a conformation change at the myosin binding sites of II, occurring during superpptn., resulting in looser binding of nucleotide.

RN 58-64-0 CAPLUS CN Adenosine 5'-(trihydrogen diphosphate) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1963:35225 CAPLUS

DN 58:35225 OREF 58:6055b-d

TI Nucleic acid synthesis in the thyroid

AU Hall, Reginald

CS Harvard Med. School, Boston, MA

SO Biochim. Biophys. Acta (1962), 61, 530-7

DT Journal

LA English

The incorporation of HC1402H into the bases of ribonucleic acid (RNA) and deoxyribonucleic acid (DNA) in slices of sheep, pig, and calf thyroid was studied. For comparison with thyroid, RNA synthesis was measured in other calf and sheep tissues. Also, labeling of adenine, guanine, and hypoxanthine was detd. Significant RNA synthesis occurred in the slices. DNA formation was much less active. In calf thyroid, purine synthesis was very rapid, and 44% of the label was incorporated into the C-8 of RNA adenine, indicating that there was significant purine formation via the full de novo pathway. 5-Amino-4-imidazole carboxamide (I) stimulated purine formation, and I ribonucleoside was even more effective. Glucose alone stimulated purine synthesis slightly, but glucose plus I was as effective as I ribonucleoside.

IT 2627-69-2, Imidazole-4-carboxamide, 5-amino-1-.beta.-D-ribofuranosyl-

(in purine formation by thyroid, D-glucose and)

RN 2627-69-2 CAPLUS

CN 1H-Imidazole-4-carboxamide, 5-amino-1-.beta.-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 23 OF 23 CAPLUS COPYRIGHT 2002 ACS

AN 1962:68377 CAPLUS

DN 56:68377

OREF 56:13239g-i

TI Biosynthesis of the purines. XXVIII. Mechanism of action of adenylosuccinase

AU Miller, Richard W.; Buchanan, John M.

CS Massachusetts Inst. of Technol., Cambridge

SO J. Biol. Chem. (1962), 237, 491-6

DT Journal

LA Unavailable

 $^{\mathrm{AB}}$ The mechanism of addn. of 5-amino-1-ribosyl-4-imidazolecarboxamide 5'-phosphate and adenosine 5'-phosphate to the double bond of fumaric acid was studied through reversal of the reactions catalyzed by adenylosuccinase in a tritiated medium. The distribution of T in the aspartate moiety of the resulting intermediate was analyzed with bacterial aspartase. Since the aspartate moiety shows the same specific distribution of T label as the product of bacterial aspartase, it must be concluded that the stereospecificity of the cleavage reaction is identical to that of the enzymic removal of NH3 from L-aspartate. Since the latter reaction and the hydration of fumarate by fumarase occur by a trans mechanism, this same general type of mechanism can now be applied to the reactions catalyzed by adenylosuccinase. The complete stereospecificity of the reaction is detd. by (a) the trans structure of fumarate, (b) the L-configuration of the product, and (c) the trans addn. of elements across the double bond of fumarate. The lack of noticeable H isotope effect on the rate of the cleaving reaction requires that H be added or removed in a step distinct from the rate-limiting process. This conclusion, in turn, requires the participation of some intermediate or complex.

IT 61-19-8, 5'-Adenylic acid 7322-81-8, Imidazole
-4-carboxamide, 5-amino-1-ribofuranosyl-, 5'-phosphate
(addn. to fumaric acid double bond by adenylosuccinic lyase)

RN 61-19-8 CAPLUS

CN 5'-Adenylic acid (8CI, 9CI) (CA INDEX NAME)

RN

7322-81-8 CAPLUS
Imidazole-4-carboxamide, 5-amino-1-ribofuranosyl-, 5'(dihydrogenphosphate) (8CI) (CA INDEX NAME) CN

Relative stereochemistry.

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